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### APPLICATION

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#### UNITED STATES LETTERS PATENT

on

# USE OF PHOSPHOROTHIOLATE POLYNUCLEOTIDES IN LIGATING NUCLEIC ACIDS

by

Alex B. Burgin, Jr.

and

Lance J. Stewart

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Attorneys

CAMPBELL AND FLORES
4370 La Jolla Village Drive, Suite 700
San Diego, California 92122

# USE OF PHOSPHOROTHIOLATE POLYNUCLEOTIDES IN LIGATING NUCLEIC ACIDS

This application claims the benefit of U.S. Provisional Application No. 60/290,313, filed 5 May 10, 2001, and is incorporated herein by reference.

This invention was made with government support under grant number GM58596-02 awarded by the National Institutes of Health. The United States Government has certain rights in this invention.

### 10 BACKGROUND OF THE INVENTION

This invention relates generally to molecular cloning and, more specifically, to ligating polynucleotides.

The techniques of molecular cloning are used in 15 virtually all realms of scientific research, from anthropology to forensic science to drug discovery. The role of molecular cloning in scientific research is expanding into new territory and increasing in importance, especially as techniques of molecular cloning are becoming easier to access and perform. The sequencing of the human 20 genome is a prominent example of the contribution of molecular cloning to both basic science and medical research. The techniques of molecular cloning will also be central in the next phase of the human genome project, 25 uncovering the functions of hundreds of newly identified human genes. This phase of the project will yield insights into the underlying causes of human disease, resulting in improved diagnostics and therapeutics. Increased efficiency in molecular cloning methods will accelerate progress in 30 this, as well as many other areas of scientific endeavor.

Ligation of polynucleotides is a key step in the process of molecular cloning. Ligation is the joining of two nucleic acid molecules by the formation of a phosphodiester bond, which is the naturally occurring 5 linkage between the nucleotides that make up a nucleic acid molecule. The ligation step in a molecular cloning procedure is typically performed using a ligase enzyme. This step is generally time-consuming because ligase enzymes can require incubation times of several hours to achieve optimal 10 reaction efficiency. In recent years, a ligation method that works without a ligase enzyme has been developed and commercialized. The method employs a sequence specific viral topoisomerase enzyme. Although generally more time efficient, the "topo cloning" method is less flexible than 15 standard cloning methods. For example, the method requires that a special sequence be incorporated into the DNA to be cloned. Further, "topo cloning" limits the user to a particular cloning site and therefore cannot be used for direct cloning of DNAs into user-selected restriction sites.

Thus, there exists a need for improved methods for ligating nucleic acids. The present invention satisfies this need and provides related advantages as well.

### SUMMARY OF THE INVENTION

The invention provides a method of nonenzymatic ligation of a nucleic acid. The method consists of contacting a polynucleotide-3' phosphorothiolate with an acceptor polynucleotide under conditions that allow formation of a phosphodiester bond between the polynucleotide-3' phosphorothiolate and the acceptor polynucleotide.

10 The invention also provides methods of molecular cloning. In one embodiment, the method consists of contacting an insert comprising a polynucleotide-3' phosphorothiolate with an acceptor vector under conditions that allow formation of a phosphodiester bond between the insert and the acceptor vector to generate a vector 15 comprising an insert polynucleotide. In another embodiment, the method of molecular cloning consists of contacting a vector comprising a polynucleotide-3' phosphorothiolate with an acceptor polynucleotide, under 20 conditions that allow formation of a phosphodiester bond between the vector and the acceptor polynucleotide to generate a vector comprising the acceptor polynucleotide.

The invention further provides a compound

25 consisting of a polynucleotide-3' phosphorothiolate and a

kit containing a polynucleotide-3' phosphorothiolate. Such

a kit can be used to ligate nucleic acids.

Also provided is a method of ligating a nucleic acid using a non-sequence specific topoisomerase. The method consists of contacting a polynucleotide-5'

phosphorothiolate with a non-sequence specific
topoisomerase, or a fragment or modification thereof, and an
acceptor polynucleotide under conditions that allow
formation of a phosphodiester bond between the

5 polynucleotide-5' phosphorothiolate and the acceptor
polynucleotide, with the proviso that the polynucleotide-5'
phosphorothiolate does not contain the nucleotide sequence
G(C/T)CCTT (SEQ ID NO:5).

The invention provides a composition containing a polynucleotide-5' phosphorothiolate and a non-sequence specific topoisomerase, and a kit containing a polynucleotide-5' phosphorothiolate and a non-sequence specific topoisomerase.

#### BRIEF DESCRIPTION OF THE DRAWINGS

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Figure 1 is a schematic representation of the reaction mechanism of polynucleotide-3' phosphorothiolate ligation to an acceptor polynucleotide.

Figure 2 is a photograph of a denaturing
20 polyacrylamide gel which shows the product of ligating a
duplex 3'-phosphorothicate polynucleotide and an acceptor
oligonucleotide.

Figure 3A shows the nucleotide sequence of each strand of duplex substrate TCRIBamS/TCRIBamAS (SEQ ID NOS:8 and 9). Figure 3B shows the nucleotide sequence of each strand of duplex substrate TCRIBstBS/TCRIBstBAS (SEQ ID NOS:10 and 11). Figure 3C shows the duplex oligonucleotide product generated by Topo65 cleavage of a TCRIBamS/TCRIBamAS duplex substrate (SEQ ID NOS:14 and 15). Figure 3D shows

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the nucleotide sequence of two cleaved TCRIBamS/TCRIBamAS oligonucleotide products ligated at a GATC overhang (SEQ ID NOS:16 and 17).

Figure 4 is a photograph of a polyacrylamide gel which shows that Topo65 forms a covalent complex with oligonucleotide substrates.

Figure 5A is a schematic representation of substrate oligonucleotide pairs that contain 5'

10 phosphorothiolate moieties. Figure 5B is a photograph of a polyacrylamide gel which shows products of Topo65-mediated ligation of 5' phosphorothiolate oligonucleotides. Figure 5C is a photograph of an denaturing polyacrylamide gel which shows products of Topo65-mediated ligation of

5' phosphorothiolate oligonucleotides.

Figure 6A is a schematic representation of a vector adapted with polynucleotide-5' phosphorothicate oligonucleotides. Figure 6B is a schematic representation of plasmids generated by Topo65-mediated recircularization of adapted linear plasmid DNA.

Figure 7A is a schematic representation of the chemical synthesis of 5'-phosphorothiolate nucleic acid 5'-S-dimethoxytrityl-3'-O-cyanoethyl-NN-diisopropyl-phosphoramidite-2'-deoxy-N-benzoyl-cytosine. Figure 7B is a schematic representation of the chemical synthesis of 5'-S-dimethoxytrityl-3'-O-cyanoethyl-NN-diisopropyl-phosphoramidite-2'-deoxy-N-isobutyryl guanosine.

Figure 8 is a schematic representation of the one-step synthesis of a polynucleotide-3' phosphorothiolate.

## DETAILED DESCRIPTION OF THE INVENTION

This invention is directed to methods of ligating a polynucleotide.

In one embodiment, the invention is directed to 5 non-enzymatic ligation of polynucleotides. Non-enzymatic ligation of polynucleotides is achieved when a chemical reaction resulting in the formation of a phosphodiester bond occurs between a 3'-phosphorothiolate derivatized polynucleotide and an acceptor nucleic acid. In comparison 10 with standard ligation methods that require an enzyme, nonenzymatic ligation provides the advantages of convenience, reduced cost and increased flexibility in designing reaction conditions. A polynucleotide-3' phosphorothiolate can be ligated to any complementary acceptor polynucleotide 15 containing a 5'-OH group. Thus, a variety of polynucleotides can be ligated using the methods of the invention. For example, a polynucleotide-3' phosphorothiolate and acceptor polynucleotide can be RNA or DNA molecules, can be single or double stranded, can be linear, circular, or branched structures, and can contain a 20 variety of functional domains, motifs, tags or moieties. The methods of the invention can be applied, for example, to generating RNA molecules, DNA molecules, RNA-DNA hybrids, vectors and inserts useful for molecular cloning, and 25 complex polynucleotide structures.

In another embodiment, the invention is directed to topoisomerase-mediated ligation between a polynucleotide 5' phosphorothiolate and an acceptor nucleic acid. The method involves trapping a non-sequence specific

topoisomerase on a polynucleotide by covalent binding of the enzyme to a 5'-phosphorothiolate moiety. A non-sequence specific topoisomerase bound to a polynucleotide 5' phosphorothiolate cleaves the polynucleotide strand at the 5'-phosphorothiolate site, generating a terminal end overhang. The topoisomerase can then ligate the bound polynucleotide to an acceptor polynucleotide having a complementary terminal end overhang.

Compared to a sequence specific topoisomerase

10 which requires a defined recognition sequence to bind to and cleave a polynucleotide, a non-sequence specific topoisomerase has the advantage of allowing to user flexibility in the nucleic acid sequences present at the terminal ends of an insert polynucleotide. This flexibility allows the user to generate topoisomerase-bound polynucleotides having terminal end overhangs of their choice. Terminal end overhangs can be selected to accommodate any restriction endonuclease-generated terminal end overhang on an acceptor polynucleotide, allowing

20 directional and non-direction ligation of an acceptor polynucleotide to a topoisomerase-bound polynucleotide.

As used herein, the term "ligation" is intended to mean the formation of a 5'-3' phosphodiester bond within a polynucleotide or between two or more polynucleotides. As used herein, the term "nonenzymatic ligation" is intended to mean ligation that occurs in the absence of an enzyme that catalyzes or facilitates the formation of a phosphodiester bond.

As used herein, the term "polynucleotide" is 30 intended to mean a chain of two or more nucleotide 5'-

monophosphate residues linked through one or more phosphodiester bonds. A nucleotide of a polynucleotide can contain a variety of glycose moieties, such as, for example, D-ribose and D-2-deoxyribose, as well as modified glycose moieties such as cytarabine. Therefore, a polynucleotide encompasses ribonucleic acid (RNA) or deoxyribonucleic acid (DNA), a hybrid of RNA and DNA, as well as RNA and DNA molecules containing nucleotides which have modified glycose moieties.

A nucleotide of a polynucleotide can contain any 10 nucleic acid base, including both naturally occurring and modified bases. Examples of naturally occurring bases include quanine, adenine, thymine, cytosine and uridine. Examples of modified bases include 4-thio-uridine, 15 pseudouridine, 2'-deoxy-uridine, 5-fluoro-uridine, 5-bromouridine, 5-iodo-uridine, 2'-amino-uridine, 2'-fluorouridine, 2'-fluoro-cytidine, 2'-amio-butyryl-pyrene-uridine, 5-fluoro-cytidine, ribo-thymidine, 5-methyl-cytidine, inosine, purine ribonucleoside, 2-aminopurine, 2,6diaminopurine, N<sup>3</sup>-methyl uridine and ribavirin. A variety 20 of other structures can be incorporated into a synthetic base. For example, a base can contain 3' and 5' modifications such as 3'-puromycin, 3'-inverted deoxy thymidine, 3'-thioate linkage, 5'-fluorescein, 5'-biotin, 25 5'-Cy3, 5'-tetrachloro-fluorescein, and other moieties, with and without atomic spacers.

A polynucleotide can be naturally occurring or synthetically produced. For example, a polynucleotide can be isolated from an organism or synthesized using various methods, such as automated methods well known in the art. A

naturally occurring polynucleotide can be, for example, an RNA such as an mRNA, a DNA such as a cDNA or genomic DNA, and can represent the sense strand, the anti-sense strand, or both.

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A polynucleotide can be a single stranded, duplex or branched polynucleotide. As used herein, the term "duplex polynucleotide" is intended to mean a polynucleotide having two strands associated together by hydrogen bonding.

- 10 A strand of a "duplex polynucleotide" can contain one or more mismatched, absent or additional nucleotides that do not associate with the cognate nucleotide in the partner strand, so long as the duplex remains associated under conditions that allow the formation of a phosphodiester bond
- between a polynucleotide-3' phosphorothiolate or a polynucleotide-5' phosphorothiolate and an acceptor polynucleotide. A duplex polynucleotide includes polynucleotides of both synthetic and natural origin. Thus, a duplex polynucleotide can be, for example, cDNA, genomic
- DNA, RNA, mRNA, synthetic DNA, including, for example, annealed complementary oligonucleotides or polynucleotides. A polynucleotide of natural origin can be derived from any eukaryotic, prokaryotic or viral source. Duplex DNA can have blunt ends, 3' terminal end overhangs and 5' terminal
- 25 end overhangs. Duplex DNA can further contain a tag or moiety, such as a tag useful for detection or purification.

As used herein, the term "polynucleotide-3' phosphorothiolate" is intended to mean a polynucleotide that contains at least one 3'-phosphorothiolate moiety.

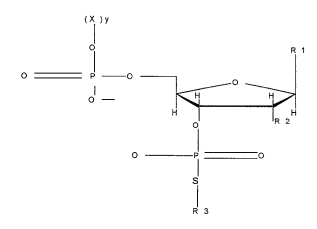
30 As used herein, the term "3'-phosphorothiolate moiety" is intended to mean a nucleotide having at least one phosphate

group linked by a phosphodiester bond at the 3' position of the sugar ring, the phosphate group having a phosphate oxygen substituted by sulfur. The sulfur can be bound to a chemical group, in particular a moiety results in the 5 formation of a good leaving group.

A polynucleotide-3' phosphorothiolate is a compound having the following structure:

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wherein,

X is a nucleotide;

Y is a positive integer;

R1 is a nucleotide base;

R2 is a hydrogen atom or hydroxyl; and

R3 is a halo, alkyl, substituted alkyl, sulfonate moiety, phenyl or substituted phenyl.

The term "alkyl" denotes such radicals as methyl, ethyl, n-propyl, isopropyl, n-butyl, sec-butyl, tert-butyl, 20 amyl, tert-amyl, hexyl and the like. A preferred alkyl group is methyl.

The term substituted alkyl groups is intended to mean alkyl groups, such as C1 to C6 and C7 to C12 alkyl groups that are substituted by one or more halogen, hydroxy, protected hydroxy, oxo, protected oxo, cyclohexyl, naphthyl, 5 amino, protected amino, (monosubstituted) amino, protected (monosubstituted) amino, (disubstituted) amino, quanidino, heterocyclic ring, substituted heterocyclic ring, imidazolyl, indolyl, pyrrolidinyl,  $C_1$  to  $C_7$  alkoxy,  $C_1$  to  $C_7$ acyl,  $C_1$  to  $C_7$  acyloxy, nitro,  $C_1$  to  $C_7$  alkyl ester, carboxy, 10 protected carboxy, carbamoyl, carboxamide, protected carboxamide, N-( $C_1$  to  $C_6$  alkyl)carboxamide, protected N-( $C_1$ to  $C_6$  alkyl)carboxamide, N,N-di( $C_1$  to  $C_6$  alkyl)carboxamide, cyano, methylsulfonylamino, thio,  $C_1$  to  $C_4$  alkylthio or  $C_1$  to  $C_4$  alkyl sulfonyl groups. The substituted alkyl groups may 15 be substituted once or more with the same or with different substituents.

Examples of the above substituted alkyl groups include the chloromethyl, bromomethyl, iodomethyl, trifluoromethyl, 6-hydroxyhexyl, 2,4-dichloro(n-butyl), 2-20 aminopropyl, chloroethyl, bromoethyl, fluoroethyl, iodoethyl, chloropropyl, bromopropyl, fluoropropyl, iodopropyl and the like.

The term "substituted phenyl" specifies a phenyl group substituted with one or more, and preferably one or two, moieties chosen from the groups consisting of halogen, hydroxy, protected hydroxy, cyano, nitro, C<sub>1</sub> to C<sub>6</sub> alkyl, C<sub>1</sub> to C<sub>7</sub> alkoxy, C<sub>1</sub> to C<sub>7</sub> acyl, C<sub>1</sub> to C<sub>7</sub> acyloxy, carboxy, protected carboxy, carboxymethyl, protected carboxymethyl, hydroxymethyl, protected hydroxymethyl, amino, protected amino, (monosubstituted) amino, protected

(monosubstituted) amino, (disubstituted) amino, carboxamide,
protected carboxamide, N-(C1 to C6 alkyl) carboxamide,
protected N-(C1 to C6 alkyl) carboxamide, N, N-di(C1 to C6
alkyl) carboxamide, trifluoromethyl, N-((C1 to C6
alkyl) sulfonyl) amino, N-(phenylsulfonyl) amino or phenyl,
substituted or unsubstituted, such that, for example, a
biphenyl results.

Examples of the term "substituted phenyl" include a mono- or di(halo)phenyl group such as 2, 3 or 4-10 chlorophenyl, 2,6-dichlorophenyl, 2,5-dichlorophenyl, 3,4dichlorophenyl, 2, 3 or 4-bromophenyl, 3,4-dibromophenyl, 3-chloro-4-fluorophenyl, 2, 3 or 4-fluorophenyl and the like; a mono or di(hydroxy)phenyl group such as 2, 3 or 4hydroxyphenyl, 2,4-dihydroxyphenyl, the protected-hydroxy 15 derivatives thereof and the like; a nitrophenyl group such as 2, 3 or 4-nitrophenyl; a cyanophenyl group, for example, 2, 3 or 4-cyanophenyl; a mono- or di(alkyl)phenyl group such as 2, 3 or 4-methylphenyl, 2,4-dimethylphenyl, 2, 3 or 4-(iso-propyl) phenyl, 2, 3 or 4-ethylphenyl, 2, 3 or 4-(npropyl)phenyl and the like; a mono or di(alkoxyl)phenyl 20 group, for example, 2,6-dimethoxyphenyl, 2, 3 or 4methoxyphenyl, 2, 3 or 4-ethoxyphenyl, 2, 3 or 4-(isopropoxy) phenyl, 2, 3 or 4-(t-butoxy) phenyl, 3-ethoxy-4methoxyphenyl and the like; 2, 3 or 4-25 trifluoromethylphenyl; a mono- or dicarboxyphenyl or (protected carboxy) phenyl group such as 2, 3 or 4carboxyphenyl or 2,4-di(protected carboxy)phenyl; a mono-or di(hydroxymethyl)phenyl or (protected hydroxymethyl)phenyl such as 2, 3, or 4-(protected hydroxymethyl)phenyl or 3,4di(hydroxymethyl)phenyl; a mono- or di(aminomethyl)phenyl or

(protected aminomethyl) phenyl such as 2, 3 or 4-

(aminomethyl)phenyl or 2,4-(protected aminomethyl)phenyl; or
a mono- or di(N-(methylsulfonylamino))phenyl such as 2, 3 or
4-(N-(methylsulfonylamino))phenyl. Also, the term
"substituted phenyl" represents disubstituted phenyl groups
wherein the substituents are different, for example, 3methyl-4-hydroxyphenyl, 3-chloro-4-hydroxyphenyl, 2-methoxy4-bromophenyl, 4-ethyl-2-hydroxyphenyl, 3-hydroxy-4nitrophenyl, 2-hydroxy 4-chlorophenyl and the like.

The terms "halo" and "halogen" refer to fluoro,

10 chloro, bromo or iodo groups. An R3 group can contain one
or more halogens, which are the same or different.

A sulfonate moiety can include, for example, a phenyl sulfonate, a substituted phenyl sulfonate, such as a halogen-substituted phenyl sulfonate, an alkyl sulfonate, such as methylsulfonate, and a substituted alkylsulfonate, such as trifluoromethylsulfonate. Sulfonate groups also include, for example, p-toluenesulfonates such as Tosyl, Brosyl, Mesyl, and Trifyl.

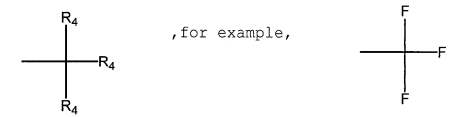
Therefore, a polynucleotide- 3'phosphorothiolate

20 of the invention can contain one of a variety of R3 groups
which can function as a leaving group in a nonenzymatic
ligation reaction. Exemplary R3 groups include the alkyl,
substituted alkyl, phenyl, substituted phenyl, and alkyl,
phenyl, substituted alkyl and substituted phenyl sulfonate

25 groups shown below:

A trisubstituted alkyl, such as a halogen substituted alkyl, wherein R4 is any halogen or combination of halogen and hydrogen,

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Trifluoromethyl.

A substituted phenyl, including a halogensubstituted phenyl, wherein R5 is any halogen, such as:

$$R_5$$

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which can be, for example, fluorophenyl:

A trisubstituted phenyl, wherein R4 is any halogen or combination of halogen and hydrogen,

$$R_4$$
 $R_4$ 

for example, trifluorophenyl,

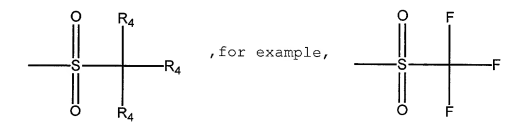
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Other exemplary substituted phenyls include, for example:

$$N=N=N$$

A trisubstituted alkylsulfonate, wherein R4 is any halogen or combination of halogen and hydrogen,



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trifluoromethyl sulfonate.

A phenyl sulfonate, for example,

such as a substituted phenyl sulfonate, wherein R5 is a halogen, including, for example,

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An exemplary substitued phenyl sulfonate is fluorophenyl sufonate:

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A polynucleotide-3' phosphorothiolate contains a phosphorothiolate moiety at the 3' carbon of the sugar ring of a nucleotide. The nucleotide is linked at its 5' carbon by a phosphodiester bond to another nucleotide, Xy. Since Y is a positive integer, the nucleotide X can be linked to any number of additional nucleotides. For example, a

polynucleotide-3' phosphorothiolate can contain as few as two nucleotides, such as when y=1, to as many as thousands of nucleotides. The nucleotides can be naturally occurring, synthetic, modified, or a combination thereof.

- As used herein the term "3'-SNP" is intended to mean a 3'-phosphorothiolate moiety in which the sulfur molecule is bound to a nitrophenyl, that is when R3 = nitrophenyl.
- As used herein the term "polynucleotide-3' phosphorothiolate precursor" is intended to mean a compound that undergoes a chemical reaction with an activator, as defined herein, to produce a polynucleotide-3' phosphorothiolate. A polynucleotide-3' phosphorothiolate precursor is a polynucleotide-3' phosphorothiolate in which the substituted sulfur molecule can undergo nucleophilic attack on an activator, displacing a molcule in the activator, and resulting in the formation of a distinct polynucleotide-3' phosphorothiolate. For example,

  20 polynucleotide-3' phosphorothiolate precursor can be a polynucleotide-3' phosphorothiolate wherein R3 is a hydrogen atom.

As used herein the term "activator" is intended to mean a compound or combination of compounds that undergoes a chemical reaction with a "polynucleotide-3' phosphorothicate precursor" to produce a polynucleotide-3' phosphorothicate. An activator contains a molecule that is displaced upon nucleophilic attack by a sulfur atom of a polynucleotide-3' phosphorothicate precursor, resulting in the formation of a covalent bond between the activator and

polynucleotide-3' phosphorothiolate precursor to produce a polynucleotide-3' phosphorothiolate. An activator therefore contains a molecule which can be displaced by a sulfur atom, such as, for example, iodine or bromine.

As used herein, the term "acceptor polynucleotide" 5 is intended to mean a polynucleotide that contains a 5'-OH group that can form a phosphodiester bond with a nucleic acid. An acceptor polynucleotide can contain one or more 5'-OH groups that can form a phosphodiester bond with a nucleic acid. For example, an acceptor nucleic acid 10 containing one 5'-OH group can ligate to one end of a nucleic acid, such as a 3'-phosphorothiolate polynucleotide or topoisomerase-bound 5'-phosphorothiolate polynucleotide. An acceptor nucleic acid containing a 5'-OH group at each 15 terminal end can ligate to two ends of a nucleic acid, such as, for example, in cloning an insert into a circular nucleic acid such as a plasmid. A branched acceptor nucleic acid can contain one or more 5'-OH groups that can ligate to one or more nucleic acids to generate, for example, larger 20 or more complexed branched nucleic acid species.

A acceptor polynucleotide can also comprise a polynucleotide 3'- or 5'-phosphorothiolate. For example, self-ligation can occur in a 3'- or 5'-polynucleotide phosphorothiolate containing an 5'-OH group that can form a phosphodiester bond.

As used herein, the term "non-sequence specific topoisomerase" refers to a class of enzymes, topoisomerase type I, found in both prokaryotes and eukaryotes which catalyze the interconversion of different topological

isomers of DNA. An exemplary non-sequence specific topoisomerase is human topoisomerase I which has the amino acid sequence referenced as SEQ ID NO:2, encoded by the nucleotide sequence referenced as SEQ ID NO:1. These

5 sequences are available in the Genbank database as accession number XM\_018038. A non-sequence specific topoisomerase can cleave multiple different nucleic acid sequences and can be specifically targeted to a nucleic acid sequence of a polynucleotide 5'-phosphorothiolate. In contrast, a

0 sequence specific topoisomerase cleaves primarily at a particular recognition sequence in a nucleic acid and does not substantially cleave non-recognition sequence containing nucleic acids.

Topoisomerases function in maintaining a specific 15 DNA topology during nuclear processes such as transcription, replication, recombination, repair, chromatin assembly and chromosome segregation by cleaving and rejoining DNA strands. Topoisomerase I enzymes bind to and cleave only one of the two DNA strands, and ligate the strand to another 20 DNA strand. Strand cleavage by type I topoisomerases occurs through transesterification of an active site tyrosine nucleophile to one strand of duplex DNA to generate a covalent DNA-(3'-phosphotyrosyl)-topoisomerase intermediate and a 5'-hydroxyl (5'-OH) DNA leaving group. 25 ligation by type I topoisomerases occurs through a second transesterification event in which a 5'-OH attacks the covalent DNA-(3'-phosphotyrosyl)-topoisomerase intermediate and displaces the topoisomerase. A "topoisomerase activity" therefore includes binding to a polynucleotide, 30 polynucleotide strand cleavage and polynucleotide strand ligation.

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The term topoisomerase encompasses native nonsequence specific topoisomerases from all species including non-sequence specific viral topoisomerases. The term "topoisomerase" also encompasses polypeptides containing 5 minor modifications of a native topoisomerase sequence, and fragments of a full-length native topoisomerase, so long as the modified polypeptide or fragment retains one or more biological activities of a native topoisomerase, such as the abilities to bind to, cleave, and ligate DNA. A 10 modification of a topoisomerase can include more additions, deletions, or substitutions of amino acids, so long as a biological activity of a native topoisomerase is retained. For example, a modification can serve to alter the stability or activity the polypeptide, or to facilitate its 15 purification.

A modified topoisomerase can contain amino acid analogs, derivatives and mimetics. Such modifications and functional equivalents of amino acids are well known to those skilled in the art. Amino acid analogs include modified forms of naturally and non-naturally occurring amino acids. Such modifications can include, for example, substitution or replacement of chemical groups and moieties on the amino acid or by derivitization of the amino acid. Amino acid mimetics include, for example, organic structures which exhibit functionally similar properties such as charge and charge spacing characteristic of the reference amino acid. For example, an organic structure which mimics arginine would have a positive charge moiety located in similar molecular space and having the same degree of mobility as the  $\epsilon\text{-amino}$  group of the side chain of the naturally occurring amino acid. Those skilled in the art

know or can determine what structures constitute functionally equivalent amino acid analogs and amino acid mimetics.

A "fragment" of a topoisomerase is intended to

5 mean a portion of a topoisomerase that retains at least
about the same activity as a native topoisomerase. A
fragment of a topoisomerase can contain a modification. The
Topo65 human topoisomerase I enzyme described herein is an
exemplary fragment of a topoisomerase that contains a

10 modification consisting of a polyhistidine tag.

The human topoisomerase I enzyme termed "Topo65" is encoded by the nucleic acid sequence referenced as SEQ ID NO:1. The Topo65 polypeptide contains amino acids

15 215 through 765 of human topoisomerase I, and is referenced as SEQ ID NO:2. Topo65 contains a series of six histidine residues incorporated at the amino terminus of the polypeptide, from amino acid residues 12 through 17.

The term "topoisomerase" includes recombinantly

20 expressed topoisomerases expressed in cells or cell lysates,
and includes chemically synthesized topoisomerases. A

topoisomerase can contain an exogenous amino acid sequence,
such as, for example, a tag that facilitates purification or
identification. Exemplary tags include histidine tags,

25 glutathione-S transferase tags, FLAG tags and myc tags.

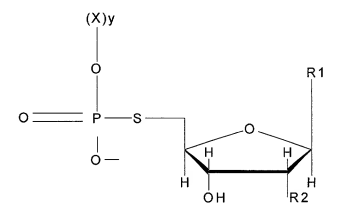
Other chemical tags such as biotin and fluorescent or
radioactive tags can be present on a topoisomerase
polypeptide or nucleic acid molecule.

The term "topoisomerase" excludes enzymes that are structurally distinct from a topoisomerase, but have enzymatic activities or reaction mechanisms similar to those of a topoisomerase. Therefore, the term "topoisomerase" specifically excludes enzymes such as integrases and recombinases, including for example,  $\lambda$  integrase, that cleave and ligate DNA but are structurally distinct from a topoisomerase of the invention.

The term "isolated", when used in reference to a topoisomerase of the invention is intended to mean a topoisomerase that is substantially removed or separated from components with which it is naturally associated.

The term "substantially the same amino acid sequence" when used in reference to a topoisomerase amino 15 acid sequence is intended to mean an amino acid sequence having at least about 70% identity with respect to a reference amino acid sequence, and retaining comparable activity characteristic of the topoisomerase defined by the reference amino acid sequence. Polypeptides having 20 "substantially the same amino acid sequence" will have at least about 80%, such as about 90% amino acid identity with respect to the reference amino acid sequence; including 95% and 98% amino acid identity with respect to the reference amino acid sequence. It is recognized, however, that polypeptides containing less than the described levels of sequence identity, arising as splice variants or that are modified by conservative amino acid substitutions are also encompassed within the scope of the present invention.

As used herein, the term "polynucleotide-5'phosphorothiolate" is intended to mean a compound having the following structure:



5 wherein,
X is a nucleotide;
Y is a positive integer;
R1 is a nucleotide base; and
R2 is a hydrogen atom or hydroxyl.

As shown, a polynucleotide-5' phosphorothiolate contains a phosphorothiolate moiety at the 5' carbon of the sugar ring of a nucleotide, forming a phosporothiolate linkage to a nucleotide, X. Since Y is a positive integer, a nucleotide can be linked to any number of additional nucleotides. For example, a polynucleotide-5' phosphorothiolate can contain as few as two nucleotides, such as when y=1, to as many as thousands of nucleotides. The nucleotides can be naturally occurring, synthetic, modified, or a combination thereof.

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The term "polynucleotide-5' phosphorothiolate" specifically excludes polynucleotide molecules consisting of certain nucleotide sequences. In particular, the term "polynucleotide-5' phosphorothiolate" specifically excludes 5 nucleic acid molecules containing the sequences G(C/T)CCTT(SEQ ID NO:5), AAA AAG ACT TAG AAA AAN NTT T, wherein N = 5bromo-deoxyuridine (SEQ ID NO:6), AAA AAG ACN NTG AAA AAN NNN T, wherein N = 5-iodo-deoxyuridine (SEQ ID NO:7), AAA AAT NNN NCN AAG TCT TTN T, wherein N is 5-bromo-deoxyuridine (SEQ ID NO:12) and AAA AAT NNN NCA AAG TCT TTT T, wherein N is 5-iodo-deoxyuridine (SEQ ID NO:13).

As used herein, the term "vector" is intended to mean a DNA of any transmissible agent, such as a plasmid or virus, for example, into which a foreign DNA sequence can be 15 spliced in order to introduce the foreign DNA into a host cell and promote its replication and transcription therein. A vector includes plasmid vectors, such as pBR322 and pUC vectors; bacteriophage vectors, such as  $\lambda gt10$  and  $\lambda EMBL4$ ; hybrid vectors, such as cosmids, phagemids, and phasmids; baculovirus and retroviral vectors; and vectors particularly useful for cloning large nucleic acid molecules, such as bacterial artificial chromosome vectors (BACs) and yeast artificial chromosome vectors (YACs). A vector can have a variety of features that render it useful for subcloning, sequencing, and expressing nucleic acids and polypeptides. For example, a vector can contain enhancer sequences, promotor sequences, transcription termination and RNA processing signals, an origin of replication, selectable marker genes and multiple cloning sites.

As used herein, the term "host cell" is intended to mean a cell that receives a polynucleotide. A host cell can be, for example, a bacterial, yeast, plant, insect, frog, fly, fish or mammalian cell that receives a polynucleotide. A variety of strains and types of host cells are included within the definition of "host cell." For example, a variety of strains of *E. coli* bacteria are routinely used for molecular cloning. Such strains include, for example, TOP10, BL21, NovaBlue and HB101.

10 As used herein, the term "terminal end overhang" is intended to mean one or more nucleotides of a single stranded portion of a duplex polynucleotide. A terminal end overhang can be present on a 3' or 5' end of a duplex polynucleotide.

15 The invention provides a method of ligating a nucleic acid. The method consists of contacting a polynucleotide-3' phosphorothiolate with an acceptor polynucleotide under conditions that allow formation of a phosphodiester bond between the polynucleotide-3'

20 phosphorothiolate and the acceptor polynucleotide.

A polynucleotide 3-phosphorothiolate and acceptor polynucleotide are ligated together in the absence of any enzyme. Therefore, the methods of the invention involving ligating a polynucleotide 3-phosphorothiolate and an acceptor polynucleotide are also referred to as methods of non-enzymatic ligation.

A polynucleotide-3'phosphorothiolate, polynucleotide-5' phosphorothiolate, and acceptor

polynucleotide can have a variety of physical and functional attributes, allowing the generation of diverse ligation products. As such, the methods of ligating a nucleic acid can be used to construct polynucleotides for a variety of 5 applications, from basic molecular cloning to building specialized polynucleotide molecules or complexes of polynucleotide molecules. For example, a polynucleotide-3'phosphorothiolate, polynucleotide-5' phosphorothiolate and acceptor polynucleotide can be single or double stranded, 10 branched, circular or linear polynucleotides. Accordingly, the methods of the invention can be used to ligate polynucleotide-3'phosphorothiolates or polynucleotide-5' phosphorothiolates and acceptor polynucleotides to produce, for example, synthetic RNAs and DNAs, RNA-DNA hybrids, 15 nucleic acids having particular structures, including highly branched structures.

The methods of the invention for ligating a nucleic acid can be advantageously applied to molecular cloning. Either or both cloning vector and insert can contain a 3'phosphorothiolate moiety that enables non-enzymatic ligation with an acceptor polynucleotide. Single inserts, as well as multiple inserts, can be cloned into a vector. Thus, the molecular cloning methods of the invention can be used for routine cloning tasks as well as more complex cloning projects, such as, for example, gene synthesis.

A polynucleotide-3' phosphorothiolate mediates non-enzymatic ligation of a nucleic acid because the sulfur-phosphorus bond is labile, and is subject to attack by a 5'-OH group of an acceptor polynucleotide. A chemical group

bound to the sulfur molecule such as, for example, an alkyl or phenyl halide, when bound to sulfur will form a good leaving group, such that the sulfur atom will be displaced, allowing the formation of a phosphodiester bond upon nucleophilic attack by a 5'-OH of an acceptor polynucleotide. As described herein, this mechanism of action was identified through detailed study of the mechanism of topoisomerase cleavage and ligation of nucleic acid suicide substrates.

ligated using the methods of the invention for non-enzymatic ligation can have a variety of chemical, structural and functional properties. Likewise a polynucleotide-5' phosphorothiolate can have a variety of chemical, structural and functional properties. For example, a polynucleotide-3' phosphorothiolate or polynucleotide-5' phosphorothiolate can be a DNA or RNA molecule, or hybrid of DNA and RNA, and can include modified glycose moieties, modified nucleic acid bases, and tags such as detection and purification tags. A hybrid of DNA can be, for example, a DNA strand bound to a complementary RNA strand, or a single polynucleotide strand containing both ribonucleic acids and deoxyribonucleic acids.

It can be desirable to prepare a hybrid

25 polynucleotide containing DNA and 2'-O-methyl RNA, for
example, to obtain enhanced nuclease resistance and
increased binding affinity. Modified glycose moieties and
nucleic acid bases can be incorporated into a
polynucleotide-3' phosphorothiolate, for example, to

30 facilitate detection or to alter the affinity of interaction

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with another molecule. Incorporating modified bases such as deoxyinosine and uridine into a DNA molecule can, for example, make duplexes with oligonucleotides more stable and can facilitate the preparation of degeneracy in primer

5 sequences. In addition RNA polynucleotides, such as oligonucleotides, made with modified bases can have higher affinities for DNA without the inherent nuclease sensitivity and instability of RNA. The uses of modified nucleic acid bases are well known in the art. Thus, methods of

10 incorporating such modifications into a polynucleotide are known and there are many commercial sources for obtaining polynucleotides containing modified bases.

Tags and moieties, such as those useful for detection and purification of a polynucleotide, include, for example, a radiolabel, a fluorochrome, a ferromagnetic substance, a luminescent tag, molecular beacon, or a detectable binding agent such as biotin. Fluorescent tags such as fluorescein dyes, rhodamine dyes, BODIPY, and Cy3 or 5 dyes can be useful, for example, in antisence applications for tracking the intracellular location of a ligated product polynucleotide, in situ hybridization, PCR product detection and a variety of DNA fragment analysis protocols well known in the art. A tag such as biotin is useful for facilitating capture of a molecule using a streptavidin-coated medium, for example. Other tags, such as a primary amine, can be incorporated into a polynucleotide to make it receptive to incorporation with another tag or label. The use of a radioactive tag is described herein, in Example III.

A polynucleotide-3' phosphorothiolate can contain one or more incorporated 3'phosphorothiolate moieties. example, a 3' phosphorothiolate moiety can be incorporated at a single 3' end of a single or double stranded 5 polynucleotide, such as when it is desired to generate a linear polynucleotide ligation product, or at two 3' ends of a double stranded polynucleotide, such as when it is desired to generate a circularized ligation product, or when a plurality of polynucleotides are being ligated together to 10 generate a linear or circular ligation product. A 3' phosphorothiolate moiety can also be incorporated into more than one 3' end of a branched polynucleotide, such as when it is desired to generate longer or more complex polynucleotide branches or extensions. Likewise, a 15 polynucleotide-5' phosphorothiolate can contain one or more incorporated 5'phosphorothiolate moieties which can be incorporated into single stranded, double stranded and branched polynucleotides.

A polynucleotide-3' phosphorothiolate can be

20 prepared, for example, by incorporating one or more 3'phosphorothiolate moieties into a polynucleotide during
synthesis of the polynucleotide, by ligating a
polynucleotide-3' phosphorothiolate onto a particular
polynucleotide, and by using one or more oligonucleotide

25 polynucleotide-3' phosphorothiolates as PCR primers for
amplifying a desired polynucleotide. When using one
polynucleotide-3' phosphorothiolate primer, a second primer
can contain a different moiety or tag, if desired. For
example, a desired nucleic acid sequence can be amplified

30 using one polynucleotide-3' phosphorothiolate primer. A

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polynucleotide-5' phosphorothiolate can be prepared in the same manner by incorporating one or more 5'phosphorothiolate moieties during synthesis or postsynthetically.

A polynucleotide-3' phosphorothiolate can also be generated from a polynucleotide-3' phosphorothiolate precursor by reacting the precursor with an activator. polynucleotide-3' phosphorothiolate precursor contains a sulfur atom in a phosphorothiolate moiety capable of nucleophilic attack of a bond in an activator molecule to 10 produce a polynucleotide-3' phosphorothiolate containing a substituted sulfur atom which acts as a good leaving group in the methods of the invention for non-enzymatic ligation of nucleic acids. For example, the sulfur atom of a polynucleotide-3' phosphorothiolate wherein R3 is a hydrogen 15 atom, can displace an iodine atom from 4-iodo-nitrobenzene to produce a polynucleotide-3' phosphorothiolate wherein R3 is nitrophenyl (see Example VII). Therefore, an activator can be a compound having an atom which can be displaced by a sulfur atom bonded to an oxygen atom within a polynucleotide 3' phosphorothiolate, such as a compound having a halo which can be displace by nucleophilic attack. The selection of an activator will determine the R3 group bound to the polynucleotide 3'-phosphorothiolate sulfur. Therefore, an activator can be selected to generate a polynucleotide 3'-25 phosphorothiolate which contains a good leaving group. described and depicted herein, R3 groups such as alkyl halides, phenyl halides, alkyl sulfonates and phenyl sulfonates can be good leaving groups for incorporation into a polynucleotide-3' phosphorothiolate. Activators which, 30 upon reaction with a polynucleotide-3' phosphorothiolate

precursor, produce polynucleotide-3' phosphorothiolates containing such R3 groups are well known to those skilled in the art. Such activators include, for example, iodonitrobenzene, iodo-trifluorobenzene and iodo-, bromo- or other derivatives of R3 moieties described herein.

It can be desirable to generate a polynucleotide-3' phosphorothiolate from a precursor, for example, when a precursor has greater stability, is more conveniently stored or packaged, or is produced at lower cost than a 10 polynucleotide-3' phosphorothiolate. A polynucleotide-3' phosphorothiolate precursor can be used to generate a polynucleotide-3' phosphorothiolate prior to addition to an acceptor nucleic acid or in the presence of an acceptor nucleic acid, if desired. Any of the ligation methods of 15 the invention that employ a polynucleotide-3' phosphorothiolate described herein can be performed with a polynucleotide-3' phosphorothiolate generated by reaction of a polynucleotide-3' phosphorothiolate precursor and an activator.

Conditions sufficient to react a polynucleotide-3' 20 phosphorothiolate precursor and an activator will vary depending on the particular precursor and activator used. Those skilled in the art will understand how to select useful precursors and activators and will know, or can 25 determine, how to obtain a particular precursor or activator. Conditions sufficient to react an exemplary polynucleotide-3' phosphorothiolate precursor, a polynucleotide-3' phosphorothioate, and an exemplary activator, 4-iodo-nitrophenyl, are described herein in Example VII.

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A variety of analytical methods and functional assays well known in the art can be employed to determine if a particular synthesis method, including generation of a polynucleotide-3' phosphorothiolate by reaction with an 5 activator, is useful for generating a desired polynucleotide-3' phosphorothiolate or polynucleotide-5' phosphorothiolate. For example, a synthetic product an be analyzed using standard methods such as NMR, mass spectroscopy and HPLC using appropriate standards, or assayed for ligation activity using various, such as those described herein.

Ligation of a polynucleotide-3' or 5' phosphorothiolate produces a product polynucleotide which does not contain a phosphorothiolate sulfur atom. 15 Therefore, a ligated product can be conveniently separated from unligated phosphorothiolate sulfur-containing polynucleotide by a variety of separation methods, if desired. For example, sufur-containing polynucleotides can be removed from non-sulfur containing polynucleotides by 20 absorption onto a chromatographic medium that binds sulfur. Exemplary sulfur-binding molecules which can be incorporated into separation media include heavy metals, such as mercury. Alternatively, various separation methods well known in the art, including methods based on isolating a polynucleotide 25 containing a purification tag, can be emloyed when it is desired to separate ligated from non-ligated polynucleotides.

Non-enzymatic ligation occurs when an acceptor polynucleotide 5'-OH displaces the sulphur contained in the modified phosphodiester linkage of a polynucleotide-3'

phosphorothiolate. Therefore, an acceptor polynucleotide contains at least one 5'-OH group. In addition to this characteristic, an acceptor polynucleotide to be ligated using the methods of ligating polynucleotides of the invention also can have a variety of chemical, structural functional properties. As described in reference to a polynucleotide-3' phosphorothiolate, an acceptor polynucleotide can be single stranded, duplex, or branched DNA, RNA or hybrid molecule, can contain a variety of modified glycose and nucleotide bases, and can contain additional useful tags or moieties.

An acceptor polynucleotide having one or more 5'-OH groups can be prepared using a variety of methods well known in the art. For example, methods such as cleavage of 15 an acceptor polynucleotide with a restriction endonuclease and dephosphorylation using phosphatases such as bovine alkaline phosphatase and calf intestinal alkaline phosphatase can be used to generate a 5'-OH containing acceptor polynucleotide. A 5'-OH containing acceptor polynucleotide can also be synthesized using automated methods well known in the art. For example, a standard nucleic acid synthesis method typically involves protecting the 5'-OH group of a nucleic acid with a dimethoxytrityl (DMT) group. Deprotection of the synthesized polynucleotide results in the generation of a 5'-OH group. For example, a DMT group can be removed from a polynucleotide by detritylation with a dichloroacetic acid/trichloroacetic acid to generate a 5'-OH group. Preparation of polynucleotides containing a 5'-OH group is routine to those of skill in the art. Analytical methods well known in the art and functional assays, such as those described herein,

can be used to determine if a polynucleotide contains a 5'-OH group.

The methods of the invention for ligating nucleic 5 acids are performed by contacting a polynucleotide-3' phosphorothiolate with an acceptor polynucleotide. polynucleotide-3' phosphorothiolate and acceptor polynucleotide can have similar or different chemical, structural and functional properties. For example, a 10 polynucleotide-3' phosphorothiolate and acceptor polynucleotide can both be RNAs, DNAs, or hybrids, or can each have different chemical properties. Both a polynucleotide-3' phosphorothiolate and acceptor polynucleotide can be single stranded, double stranded or 15 branched polynucleotides, or alternatively, can each have different structural properties. A polynucleotide-3' phosphorothiolate and acceptor polynucleotide can have similar or different functional properties. For example, both can be vectors or inserts, each having the same or 20 different constituent functional features, such as promotors, enhancers, genes, and the like, or each can have different functional properties. Similarly, a polynucleotide-5' phosphorothiolate and acceptor nucleic acid can have similar or different chemical, structural and 25 functional activities.

A duplex acceptor polynucleic can undergo nonenzymatic ligation so long as it is sufficiently complementary to base pair with a polynucleotide-3' phosphorothiolate or polynucleotide-5' phosphorothiolate.

30 For example, polynucleotides having blunt ends, or

complementary terminal end overhangs can undergo nonenzymatic ligation.

It can be advantageous to have complete complementarity between the terminal end overhangs of a 5 polynucleotide-3' phosphorothiolate or polynucleotide-5' phosphorothiolate and an acceptor polynucleotide such that ligation occurs substantially between specific nucleic acids to allow predictable ligation products. However, having missing, extra or mismatched bases present in one terminal 10 end overhang, does not preclude the generation of a replicatable plasmid in a host cell. For example, an acceptor polynucleotide that is ligated to a polynucleotide-3' phosphorothiolate to form a plasmid can be ligated to an insert only on the two 5' ends, only on the two 3' ends or 15 on one 5' and one 3' end of the insert, transformation into a host cell can produce an intact vector due to repair of the plasmid within the host cell. Therefore, transformation of a ligated polynucleotide product into a host cell is useful for generating a polynucleotide product. 20 Transformation of ligated polynucleotides into a host cell

can optionally be used in any of the methods of the invention for ligating nucleic acids.

A variety of conditions are suitable for ligating a nucleic acid using the methods of the invention involving non-enzymatic ligation and topoisomerase-mediated ligation. Such conditions can include time of reaction, temperature of reaction, components in a reaction mixture and the particular concentration of each component in a reaction mixture. Non-enzymatic and topoisomerase-mediated ligation can occur over a range of reaction times, from several

seconds, to several minutes, to several hours. The length of time required to generate ligated products will be influenced by the temperature and reaction mixture components. For example, the generation of ligated products 5 will occur in a shorter time period when performed at a warm temperature, such as 37°C, compared to a cool temperature, such as 16°C. Times of incubation can therefore be about less that one hour, such as about one minute, about two to ten minutes, about 30 minutes to one hour, or more than one hour, including several hours, such as overnight. enzymatic and topoisomerase-mediated ligation can occur over a range of temperatures. For example, using non-enzymatic ligation, ligated products can be produced at temperatures of between -20 to  $-5^{\circ}$ C, -4 to  $3^{\circ}$ C, 4 to  $15^{\circ}$ C, 16 to  $25^{\circ}$ C, 15 26 to  $36^{\circ}$ c, 37 to  $42^{\circ}$ C, 42 to  $50^{\circ}$ C, 51 to  $60^{\circ}$ C, 60 to  $75^{\circ}$ C and 76 to 100°C. For topoisomerase-mediated ligation, ligated products can be produced at temperatures of between 4 to  $16^{\circ}$ C, 16 to  $25^{\circ}$ C, 26 to  $36^{\circ}$ C, 37 to  $42^{\circ}$ C and 42 to  $50^{\circ}$ C.

The methods of the invention for ligating polynucleotides can be performed in the presence of various chemical components. Exemplary chemical components include, for example, buffers such as Tris-HCL, Tris-acetate, potassium-acetate, magnesium acetate, Bis Tris propane-HCl, and potassium phosphate, salts such as NaCl, KCl, and MgCl2, proteins such as bovine serum albumin and IgG, detergents such as Triton X-100, CHAPS and SDS, metal chelators such as EDTA and EGTA, condensing reagents such as hexamminecobalt chloride, crowding reagents such as polyethylene glycol, oxidizing or reducing agents such as dithiothrietol and  $\beta$ - mercaptoethanol, non-reactive nucleic acids molecules such as tRNA, and other chemical components such as glycerol.

One skilled in the art can readily determine a condition for that allows formation of a phosphodiester bond between a polynucleotide-3' phosphorothiolate or a polynucleotide-5' phosphorothiolate and an acceptor polynucleotide by performing the ligation reaction under a variety of time, temperature, and buffer composition conditions followed by detection of a ligated product. A ligation product can be detected by a variety of methods well known in the art, such as the SDS-PAGE methods described herein.

The invention provides a method of molecular cloning. The method consists of contacting an insert comprising a polynucleotide-3' phosphorothicate with an acceptor vector under conditions that allow formation of a phosphodiester bond between the insert and the acceptor vector to generate a vector comprising an insert polynucleotide.

A polynucleotide-3' phosphorothiolate, such as an insert comprising a polynucleotide-3' phosphorothiolate can have various chemical, structural and functional properties, as described above. An insert can be of any size with respect to a recipient vector. For example, an insert can be a polynucleotide-3' phosphorothiolate having a few, tens, or hundreds of nucleic acids.

An insert comprising a polynucleotide-3'

25 phosphorothicate to be ligated to an acceptor polynucleotide can be blunt-ended or have terminal end overhangs complementary to a recipient vector. Selection of inserts and vectors having compatible or complementary terminal end overhangs is routine in the art, and is

described, for example in and Russell, <u>Molecular Cloning: A Laboratory Manual</u>, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York (2001), which is incorporated herein by reference. In addition, it is well known in the art that ligation between certain non-complementary terminal ends can occur within a host cell. For example, a host such as *E. coli* can repair nicks in circular plasmids, fill in missing bases, and can remove an extra bases, such as those of a terminal end overhang, *in vivo* to produce a repaired, replicatable plasmid. Accordingly, transformation of a vector containing a partially ligated insert, such as an insert ligated only at 3' ends of the insert or vector, can result in *in vivo* repair and generation of a replicatable vector in a host cell.

An insert comprising a polynucleotide-3' 15 phosphorothiolate can be prepared using a variety of methods, as described in relation to a polynucleotide-3' phosphorothiolate. For example, a polynucleotide-3' phosphorothiolate, such as an insert comprising a polynucleotide-3' phosphorothiolate, can be prepared by 20 synthesing a polynucleotide portion using standard polynucleotide synthesis methods, and then chemically modifying the polynucleotide to incorporate a 3'phosphorothiolate moiety. Alternatively, a 3'-25 phosphorothiolate moiety can be introduced into an insert by ligation, or an insert can be prepared by PCR using an oligonucleotide pair in which at least one oligonucleotide contains a 3'-phosphorothiolate moiety. A double-stranded insert comprising a polynucleotide-3' phosphorothiolate can be made by annealing a complementary polynucleotide strand 30 that can optionally contain a 3'-phosphorothiolate moiety.

Such methods can similarly be employed to prepare a polynucleotide-5'phosphorothiolate insert.

A duplex insert can be generated by annealing two complementary synthetic polynucleotides, one or both of

which can be a polynucleotide-3' phosphorothiolate. Methods for annealing nucleic acids are well known in the art and an exemplary condition for annealing two oligonucleotides is described herein.

As described above in reference to a polynucleotide-3' phosphorothiolate, an insert polynucleotide-3' phosphorothiolate can likewise contain one or more 3' phosphorothiolate moieties. An acceptor vector to be ligated to an insert polynucleotide-3' phosphorothiolate also can also have chemical, structural 15 and functional properties as described above for an acceptor polynucleotide. An acceptor vector, for example, can contain one or more 5-OH groups. A vector having one 5'-OH can be ligated to an insert to generate a linear vector comprising the insert. For duplex circular vector, such as 20 a plasmid vector, it is advantageous to have a 5'-OH at each terminal end of the vector to generate an intact plasmid in a host cell, as described above. Ligation of the 3' ends of an acceptor vector can be obtained, for example, by incorporation of 3'-phosphorothiolate moieties at one or 25 more 3' ends. Ligation can thereby occur between 5'-OH ends of the 3'-phosphorothiolate insert and the 3' ends of the vector. Such inserts and vectors containing both 5'-OH groups and 3'-phosphorothiolate moieties are designed to prevent or minimize self-ligation reactions, for example, by making 3' and 5' ends non-complementary.

The invention also provides a method of molecular cloning in which a vector contains a polynucleotide-3' phosphorothiolate. The method consists of contacting a vector containing a polynucleotide-3' phosphorothiolate with an acceptor polynucleotide, under conditions that allow formation of a phosphodiester bond between the vector and the acceptor polynucleotide to generate a vector containing the acceptor polynucleotide.

As described herein, in reference to a 10 polynucleotide-3' phosphorothiolate, a vector containing a polynucleotide-3' phosphorothiolate can have one or more 3'phosphorothiolate moieties incorporated at one or more 3'-Therefore, one or more phosphodiester bonds can or 5'-ends. be formed between a vector and an insert. For a plasmid 15 vector, is advantageous to incorporate a 3'-phosphorothiolate moiety at each terminal end, for example, at each 3' terminal end. An insert acceptor having a 5'-OH at each terminal end can ligate to the plasmid vector to produce an insert-containing vector. 20 insert-containing vector can therefore contain two or more nicks, one at each junction between the 5'-termini of the vector and 3'-termini of the insert. The insert-containing vector, when transformed into a host cell, such as a strain of E. coli, will be repaired to generate a replicatable 25 plasmid.

A vector containing a polynucleotide-3' phosphorothicate can contain nucleic acids from various organism including, for example, bacteria, bacteriophage, yeast, plants and mammals, as well as nucleic acids from viruses. A vector can contain a variety of features useful

for replication, gene expression, cloning and protein expression. For example, promotor, enhancer, poly A sequences, other regulatory sequences, genes encoding various polypeptides and selectable markers can be contained in a vector. A vector containing a polynucleotide-3' phosphorothicate can be prepared from a variety of vectors well known in the art, such as by incorporation of a 3' phosphorothicate moiety post-synthetically or during the synthesis of the vector.

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The invention provides a polynucleotide 3'phosphorothiolates, preferably in kit form. A kit contains (a) a polynucleotide-3' phosphorothiolate; and (b) a buffer in an aqueous solution in a suitable packaging 15 material. A polynucleotide 3'-phosphorothiolate can be generated by contacting a polynucleotide 3'phosphorothiolate precursor with an activator. Therefore, the invention also provides a kit containing a polynucleotide 3'-phosphorothiolate precursor and an activator. A kit of the invention is useful for ligating 20 nucleic acids. A kit includes at least one invention polynucleotide 3'-phosphorothiolate, as a separately packaged chemical reagent(s) in an amount sufficient at least one ligation. As described herein, a polynucleotide 3'-phosphorothiolate can be contained in a variety of 25 nucleic acids, including, for example, a DNA such as a cDNA or genomic DNA, an RNA, such as a mRNA or catalytic RNA, a vector such as a cloning or expression vector, viral DNA or BAC, an oligonucleotide, an insert, a branched

polynucleotide, and the like.

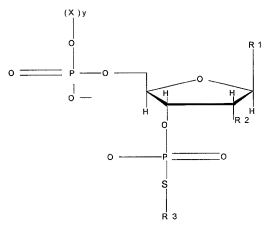
For a kit containing oligonucleotide polynucleotide 3'-phosphorothiolates, the kit will generally contain two or more oligonucleotide polynucleotide 3'-When the kit is to be used for phosphorothiolates. 5 generating a PCR product containing a 3'-phosphorothiolate, the kit will contain at least two oligonucleotides, at least one of which is a polynucleotide 3'-phosphorothiolate, that can serve as primers for PCR. Those of skill in the art can readily incorporate invention oligonucleotides in 10 combination with the provided buffer(s) and appropriate solutions for the practice of the invention methods as described herein. A kit containing a polynucleotide 3'phosphorothiolate contains a buffer that provides the proper conditions for performing a ligation, and can contain 15 control samples known to undergo ligation under appropriate conditions.

example, a polynucleotide 3'-phosphorothiolate, such as an oligonucleotide or vector polynucleotide 3'
20 phosphorothiolate and buffer in aqueous solution, are contained in packaging material, preferably to provide a sterile, contaminant-free environment. In addition, the packaging material contains instructions indicating how the materials within the kit can be employed to ligate nucleic acids. The instructions for use typically include a tangible expression describing the reagent concentration or at least one ligation method parameter, such as a time period or temperature of incubation, and the like.

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The invention provides a compound of the following formula:



wherein,

X is a nucleotide;

Y is a positive integer;

R1 is a nucleotide base;

R2 is a hydrogen atom or hydroxyl; and

R3 is a halo, alkyl, substituted alkyl, sulfonate

moiety, phenyl, substituted phenyl.

The polynucleotide-3' phosphorothiolate compound of the invention can contain a variety of substituents that function as good leaving groups at the R3 position.

Substituents that function as good leaving groups are well known to those skilled in the art, and include, for example, 15 alkyl halides, phenyl halides, alkyl sulfonates and phenyl sulfonates.

The polynucleotide-3' phosphorothiolate compound of the invention can be conveniently prepared as described in Example VII, by modifications of the method that produce 20 the polynucleotide-3' phosphorothiolate compound of the

invention, and by other methods which can be determined by those skilled in the art. Such modifications of the procedure for preparing a polynucleotide-3'phosphorothiolate can include, for example, the use of alternate solvents, buffers and temperature conditions.

The invention provides another method of ligating a nucleic acid. The method consists of contacting a polynucleotide-5' phosphorothiolate with a non-sequence specific topoisomerase, or a fragment or modification

10 thereof, and an acceptor polynucleotide under conditions that allow formation of a phosphodiester bond between the polynucleotide-5' phosphorothiolate and the acceptor polynucleotide, with the proviso that the polynucleotide-5' phosphorothiolate does not contain the nucleotide sequence

15 G(C/T)CCTT (SEQ ID NO:5).

Like the methods of the invention for nonenzymatic ligation, the methods for topoisomerase-mediated
ligation using a polynucleotide-5' phosphorothicalte can be
applied to a variety of applications, from molecular cloning
to constructing polynucleotides useful for a variety of
purposes, such as for detecting an analyte, producing
template nucleic acids, generating catalytic molecules such
as RNA molecules and RNA-DNA hybrids.

The methods for topoisomerase-mediated ligation

25 using a polynucleotide-5' phosphorothiolate involve
covalently trapping a topoisomerase at a 5'phosphorothiolate moiety of a polynucleotide-5'
phosphorothiolate. The trapped topoisomerase enzyme
catalyzes the ligation of the polynucleotide-5'

phosphorothiolate to any complementary acceptor nucleic acid having a 5'-OH group. As such, a topoisomerase can be covalently bound to a vector or insert to promote ligation to a complementary acceptor nucleic acid within seconds or minutes. A topoisomerase-bound vector can be used to conveniently and efficiently ligate to one, several or many different inserts. Such a vector can be used, for example, for routine cloning of single inserts as well as for constructing polynucleotide libraries. Conversely, a topoisomerase-bound insert can be used to conveniently and efficiently ligate to one, several, or many vectors. Such an insert could be cloned into a panel of vectors, for example, when it is desired to test multiple vectors having different functional properties for a particular application, such as gene or protein expression.

Topoisomerase I enzymes are capable of cleaving duplex polynucleotides at sites that contain a 5' bridging phosphorothiolate moiety and have been used as suicide substrates for recombinases and topoisomerases (Burgin et al. Nucleic Acids Research, 23:15, 2973-2979, (1995); Burgin et al. <u>Current Biology</u>, 5:1312-1321, (1995); Redinbo et al. Science, 279:1504-1513 (1998)). Upon cleavage of such a site, topoisomerase I becomes covalently attached to the 3' end of the broken strand through a phosphotyrosine bond between a catalytic tyrosine in the active site of topoisomerase I and the 3' end of the broken strand. generates a 5' sulfhydryl leaving strand instead of a 5'-OH, which is not a sufficient nucleophile to reverse the 30 cleavage reaction. Thus, cleavage by topoisomerase I at a 5'-bridging phosphorothiolate is irreversible in the absence of an acceptor polynucleotide. The methods of the invention employ 5'-bridging phosphorothicaltes to trap topoisomerase I in a covalent complex with a polynucleotide to generate a complex that can perform a ligation reaction with an acceptor polynucleotide having a 5'-OH.

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In the present invention, polynucleotide 5'phosphorothiolates are now used for promoting topoisomerasemediated ligation of nucleic acids. A non-sequence specific
topoisomerase is targeted to a specific nucleic acid
sequence by incorporating a 5'-bridging phosphorothiolate at
a desired site within the sequence. A non-sequence specific
topoisomerase can promote ligation between the
topoisomerase-bound polynucleotide and an acceptor
polynucleotide. The topoisomerase first cleavages at the
5'-phosphorothiolate moiety in the scissile (top) strand of
a duplex polynucleotide. The topoisomerase then catalyzes
the attack of the 5'-OH of the acceptor nucleic acid on the
topoisomerase-bound polynucleotide 5-phosphorothiolate to
form a phosphodiester bond, displacing the enzyme.

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A topoisomerase can cleave at a 5'
phosphorothiolate moiety at any location within a scissile
polynucleotide strand, allowing that the 5'
phosphorothiolate moiety is flanked by at least several

25 bases on the 5' side, and at least one base on the 3' side.
Cleavage at a 5' phosphorothiolate moiety by a topoisomerase
will generate a single stranded fragment if the cleavage
site is near the terminal end of a linear substrate and
produce a 5' overhang on the bottom strand. The single

30 stranded fragment, if relatively small, such as about 20 or
fewer bases, 10 or fewer bases, 6 or fewer bases, 4 or fewer
bases, or 2 or fewer bases, will dissociate from the duplex

to produce the overhang. A larger fragment that is not spontaneously dissociated from a duplex can be dissociated, for example, by heating to 80°C, followed by quick cooling and separation of the fragment, such as by gel purification, affinity isolation or other methods well known in the art. Therefore, an overhang generated by cleavage by a topoisomerase can contain contain few bases or many bases, depending on the site to which the topoisomerase is targeted.

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Therefore, a user of the methods of the invention for topoisomerase-mediated ligation of nucleic acids can design polynucleotide 5' phosphorothiolates having a diverse array of different overhangs. The variety of different overhangs are useful for ligating diverse acceptor polynucleotides and for cloning using desired restriction sites, and particularly for uni-directional cloning, in which an insert is ligated into a vector in a specific orientation, as dictated by the complementarity of 3'- and 5'-terminal ends of the insert and vector.

A non-sequence specific topoisomerase can be, for example, a eukaryotic or viral topoisomerase, such as a human topoisomerase I. An exemplary human topoisomerase termed Topo65 is provided herein. Topo65 is a topoisomerase fragment having topoisomerase activity, that provides several advantages over a native topoisomerase. For example, Topo65 can be conveniently purified in large quantity by metal ion affinity chromatography, it expressed in higher amount than native topoisomerase, for example, in insect cells, and is comparatively more stable than native topoisomerase.

The invention provides an isolated non-sequence specific topoisomerase, or a fragment or modification thereof having topoisomerase activity. An isolated topoisomerase can be prepared by a variety of methods wellknown in the art, for example, by recombinant expression systems described herein, and biochemical methods of purification such as precipitation and chromatographic methods. A non-sequence specific topoisomerase of the invention can be prepared, for example, by recombinant 10 expression in any compatible vector/host system. A variety of protein expression systems are well known in the art and include, for example, prokaryotic and eukaryotic expression systems. Prokaryotic expression systems are advantageous due to their ease in manipulation, low complexity growth 15 media, low cost of growth media, rapid growth rates and relatively high yields. Well known prokaryotic expression systems include, for example, E. coli bacterial expression systems based on bacteriophage T7 RNA polymerase, the trc promotor, the araB promotor and bacillus expression. Eukaryotic expression systems are advantageous because expressed polypeptides can contain eukaryotic posttranslational modifications such as O-linked glycosylation, phosphorylation and acetylation and can have improved protein folding. Well known eukaryotic expression systems 25 include, for example, expression in yeast, such as Pichia pastoris and Pichia methanolica, expression in insect systems such as the Drosophila S2 system and baculovirus expression systems and expression in mammalian cells using adenoviral vectors and cytomegalovirus promotor-containing vectors. Described herein is the purification of Topo65 30

from sf9 insect cells infected with baculovirus encoding

Topo65 (see Example IV).

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A non-sequence specific topoisomerase of the invention can be purified using a variety of methods of protein purification well known in the art. Biochemical purification can include, for example, steps such as 5 solubilization of the topoisomerase-expressing cell, isolation of the desired subcellular fractions, chromatography, such as ion exchange, size, or affinitybased chromatographies, electrophoresis, and immunoaffinity procedures. Other well-known methods are described in Deutscher et al., Guide to Protein Purification: Methods in Enzymology Vol. 182, (Academic Press, (1990)). The methods and conditions for biochemical purification of a polypeptide of the invention can be chosen by those skilled in the art, and the purification monitored, for example, by staining 15 SDS-PAGE gels containing protein samples, by immunodetection methods such as Western blotting and ELISA, and by functional assay of topoisomerase activity.

A topoisomerase can be modified, for example, to increase polypeptide stability, alter a topoisomerase activity, facilitate detection or purification, or render the enzyme better suited for a particular application, such as by altering substrate specificity. Computer programs known in the art can be used to determine which amino acid residues of a topoisomerase can be modified as described above without abolishing a topoisomerase activity (see, for example, Eroshkin et al., <a href="Comput.Appl.Biosci.">Comput. Appl. Biosci.</a> 9:491-497 (1993)). In addition, structural and sequence information can be used to determine the amino acid residues important for topoisomerase activity. For example, a topoisomerase crystal structure (Redinbo et al., supra) and comparisons of

topoisomerase amino acid sequences, such as that shown for poxvirus topoisomerases in Krogh et al. (Virology, 264, 441-451 (1999)) can provide guidance in determining amino acid residues that can be altered without abolishing topoisomerase activity.

The invention provides another kit. The kit consists of (a) a polynucleotide-5' phosphorothiolate, with the proviso that the polynucleotide-5' phosphorothiolate does not contain a nucleotide sequence selected from the group of SEQ ID NO:5, SEQ ID NO:6 or SEQ ID NO:7; (b) a non-sequence specific topoisomerase, or fragment or modification thereof having topoisomerase activity; and (c) a buffer in an aqueous solution.

A kit of the invention is useful for generating a 15 covalent complex of Topo65 and a polynucleotide and for ligating nucleic acids. A kit includes at least one invention polynucleotide-5' phosphorothiolate, with the proviso that the polynucleotide-5' phosphorothiolate does not contain a nucleotide sequence selected from the group of SEQ ID NO:5, SEQ ID NO:6 or SEQ ID NO:7 and a non-sequence specific topoisomerase of the invention as a separately packaged chemical reagent(s) in an amount sufficient at least one ligation. For a kit containing oligonucleotide 25 polynucleotide 5'-phosphorothiolates, the kit will generally contain two or more oligonucleotide polynucleotide 5'phosphorothiolates. When the kit is to be used for generating a PCR product containing a 5'-phosphorothiolate, the kit will contain at least two oligonucleotide polynucleotide 5'-phosphorothiolates that can serve as 30 primers for PCR. Oligonucleotide polynucleotide 5'-

phosphorothiolates of the kit can contain a variety of useful sequences. For example, an oligonucleotide can contain a restriction site and the sequence of a vector from which a polynucleotide is amplified. Those of skill in the art can readily incorporate invention oligonucleotides in combination with the provided non-sequence specific topoisomerase and buffer(s) and other appropriate solutions for the practice of the invention methods as described herein.

A polynucleotide 5'-phosphorothiolate can have a 10 variety of chemical, structural and function properties, as desribed above. A polynucleotide 5'-phosphorothiolate can be prepared using a variety of methods, also described Thus, a variety of polynucleotide 5'-15 phosphorothiolates can be provided in a kit of the invention. An exemplary polynucleotide 5'-phosphorothiolate that can be contained in a kit is an oligonucleotide containing a 5'-phosphorothiolate moiety. The oligonucleotide could be used with another oligonucleotide 20 to amplify a PCR product, such as a vector or insert, containing a 5'-phosphorothiolate. A polynucleotide 5'phosphorothiolate PCR product can then be covalently bound to the kit-provided Topo65 to generate a topoisomerasepolynucleotide 5'-phosphorothiolate complex capable of 25 ligating to a complementary acceptor polynucleotides. A 5'phosphorothiolate oligonucleotide can also be used, for example, to amplify a plurality of different polynucleotides. The resulting population of PCR products can then be bound to a topoisomerase and ligated, for

example, to an acceptor nucleic acid, such as a vector.

convenience, a kit can also contain vectors that can

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optionally be used with the Topo65 and polynucleotide 5'-phosphorothiolates of the invention.

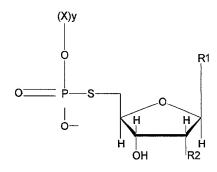
An acceptor vector or 5'-phosphorothiolate-5 containing vector can contain a variety of terminal ends, including for example, 5' or 3' terminal overhangs having nucleic acid sequences complementary to terminal end overhangs generated by a restriction endonuclease, such as a HindIII, EcoRI, BamHI or BstBI overhang. A vector bound to 10 a Topo65 through a 5' phosphorothiolate moiety, or capable of binding Topo65 through a 5'-phosphorothiolate moiety, can contain two like or different terminal overhangs. different overhangs can correspond to overhangs generated by two restriction enzymes. A topoisomerase-bound vector 15 containing overhangs corresponding to two different restriction enzyme overhangs can conveniently be used for rapid cloning of inserts having the corresponding overhangs. For example, a topoisomerase-bound vector can contain BamHI and EcoRI overhangs for cloning insert acceptor polynucleotides generated by cleavage with BamHI and EcoRI. 20

A kit containing a polynucleotide 5'phosphorothiolate and Topo65 enzyme also contains a buffer
that provides the proper conditions for performing a nonsequence specific topoisomerase-mediated ligation, and can
contain control samples known to undergo ligation under
appropriate conditions.

The invention further provides a composition containing (a) a polynucleotide-5' phosphorothiolate, with the proviso that the polynucleotide-5' phosphorothiolate does not contain a nucleotide sequence selected from the

group of SEQ ID NO:5, SEQ ID NO:6 or SEQ ID NO:7; and (b) a non-sequence specific topoisomerase, or fragment or modification thereof having topoisomerase activity.

The invention also provides a compound of the 5 formula:



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wherein,

X is a nucleotide;

Y is a positive integer;

R1 is cytosine or guanine; and

R2 is a hydrogen atom or hydroxyl.

The polynucleotide 5'-phosphorothiolate compound of the invention, such as a 5'-S-dimethoxytrityl-3'-O
cyanoethyl-NN-diisopropyl-phosphoramidite-2'-deoxy-Nbenzoyl-cytosine or 5'-S-dimethoxytrityl-3'-O-cyanoethyl-NNdiisopropyl-phosphoramidite-2'-deoxy-N-isobutyryl guanosine
can be conveniently prepared as described in Example VI and
Figure 7, by modifications of the method that produce the

polynucleotide-3' phosphorothiolate compound of the
invention, and by other methods which can be determined by
those skilled in the art. Other 5'-phosphorothiolate

compounds useful in the methods of the invention, such as those based on uridine and thymidine can be prepared, for example, as described in Mag et al., <u>Nucleic Acids Res.</u>, 19(7):1437-41 (1991).

The methods of the invention involve ligating polynucleotides, such as DNA and RNA. Both DNA and RNA molecules can be obtained using a variety of methods well known in the art. These molecules can be obtained by purification from a natural source, such as a cell from an organism, by purification from a host cell, such as a cell which overexpresses a plasmid DNA, and by chemical synthesis, such as automated oligonucleotide synthesis methods well known in the art.

The methods of the invention involve transforming a vector into a host cell. Methods for introducing a vector into a host cell are well known in the art and include, for example, various methods of transformation such as calcium chloride and electroporation and transfection such as calcium phosphate, DEAE-dextran and liposome mediated methods, viral transduction, electroporation.

The methods of the invention involve preparing duplex polynucleotides such as duplex oligonucleotides. A variety of methods for preparation of duplex polynucleotides are well known in the art. An exemplary method by which duplex oligonucleotides and polynucleotides can be prepared is by mixing equimolar concentrations of complementary strands, each at a final concentration of about 0.1 mM, in the presence of a salt, such as 6 M NaCl, and then heating

1.0

to 80°C followed by cooling to room temperature over a period of 8 hours.

It is understood that modifications which do not substantially affect the activity of the various embodiments of this invention are also included within the definition of the invention provided herein. Accordingly, the following examples are intended to illustrate but not limit the present invention.

### EXAMPLE I

### Nonenzymatic Ligation of Nucleic Acid

This example shows that a polynucleotide 3'-phosphorothicate can be ligated to an acceptor polynucleotide.

A polynucleotide 3-phosphorothiolate mediates non-15 enzymatic ligation through a mechanism analogous to that of a topoisomerase. Topoisomerase-mediated ligation is effective because the 3'-phosphotyrosine is a relatively high energy intermediate and therefore labile; formation of 20 a 5'-3' DNA phosphodiester bond is extremely stable and energetically favored. Topoisomerases require no external energy source (e.g. ATP) to catalyze the cleavage and ligation reactions. It was hypothesized that other chemically modified 3'-phosphodiesters could mimic this 25 labile intermediate. To test this hypothesis an oligonucleotide containing a 3'-S-(paranitrophenyl) moiety (termed 3'-SNP) was prepared. This compound contains a substitution of a sulfur for an oxygen because sulfurphosphorous bonds are significantly more labile than oxygen15

phosphorous bonds and therefore provides a better leaving In addition, the pKa of a thiol is three orders of magnitude lower than an alcohol, and the 4-nitro functional group reduces the pKa of the thiol even further, creating a 5 very labile 3-phosphorothiol ester. The 3'-SNP was predicted to be a good leaving group during a ligation reaction in which the acceptor polynucleotide supplies the 5'-OH necessary to displace the 4-nitrothiobenzene. reaction is depicted in Figure 1. Other thioesters and labile phophoesters can function as leaving groups in the non-enzymatic ligation reactions described herein. As shown below, a 3'-SNP within a polynucleotide served as a good leaving group, allowing a polynucleotide containing a 3'-SNP to undergo non-enzymatic ligation with an acceptor nucleic acid.

To determine if a polynucleotide having a 3'-SNP moiety could act as a leaving group during a ligation reaction and form a phosphodiester bond with another nucleic acid molecule in the absence of any enzyme, a 3'-SNP-moiety 20 was incorporated at a nick in a duplex polynucleotide. 3'-SNP-containing polynucleotide was radioactively labeled at the 5' end and was incubated for 3 hours in the presence of an acceptor oligonucleotide. The reaction mixtures were then separated on a denaturing polyacrylamide gel. Figure 2 25 is an autoradiograph which shows the products generated when a 3'-SNP-containing duplex DNA is incubated with a 20-mer acceptor oligonucleotide (lane 1), a mixture of 20-mer and 24-mer acceptor oligonucleotides (lane 2) or in the absence of acceptor oligonucleotide (lane 3). The expected ligation 30 products were observed, demonstrating that ligation can occur in the absence of any enzymes.

### EXAMPLE II

# Covalent Bond Formation Between Topo65 and 5'-Phosphorothiolate-Containing Polynucleotides

This example shows that Topo65 can cleave a duplex 5 DNA molecule containing a 5'-phosphorothiolate moiety located two or four base pairs upstream from a blunt end.

To determine if Topo65, a recombinant form of human topoisomerase I, described in detail below, can cleave near the end of a duplex DNA molecule, Topo65 was incubated

10 with oligonucleotide substrates containing 5'phosphorothiolate moieties located two or four base pairs upstream from a blunt end. The duplex substrates contained nucleic acid sequences at their 3' end that, when cleaved by Topo65, would generate 2 and 4 base overhangs compatible

15 with terminal end overhangs generated by the restriction enzymes BstBI and BamHI, respectively. The substrates contained a 5'-EcoR1 overhang on the other end of the duplex oligonucleotide so that the duplex oligonucleotides could be ligated, using standard T4 DNA ligase, onto linear plasmid

20 molecules.

Figures 3A and 3Bshows duplex substrates containing a 5'-bridging phosphorothicate "OPS" moiety located 4 or 2 base pairs from the 3' ends of a scissile strand (top strand) which is annealed to a complementary strand (bottom strand) having a 5' dimethoxytrityl group (DMT) to form a duplex having a 5' EcoRI overhang (left ends). Topo65 cleavage of the duplex results in Topo65 being trapped at the OPS moiety topo cleavage site. The

site of topoisomerase trapping is depicted by an arrow in Figures 3A and 3B. Subsequent cleavage of the scissile strand of the duplex to which Topo65 is covalently bound generates a 4 or 2 base 5' overhang which is complementary to the 5' overhangs generated by restriction digestion with the enzymes BamH1 (4 base overhang) or BstBI (2 base overhang).

Figure 3A shows the nucleotide sequences of the oligonucleotides contained in duplex oligonucleotide pair TCRIBamS/AS. The nucleotide sequences are as follows: TCRIBamS: 5' AATTCGCGGCCGCAAAAAGACTTGATC 3' (SEQ ID NO:8) and TCRIBamAS: 5' GATCAAGTCTTTTTGCGGCCGCG (SEQ ID NO:9). Figure 3B shows the nucleotide sequences of the oligonucleotides contained in duplex oligonucleotide pair TCRBstBS/AS. The nucleotide sequences are as follows: TCRBstBS: 5' AATTCGCGGCCGCAAAAAGACTTCG (SEQ ID NO:10) and TCRBstBAS: 5'CGAAGTCTTTTTGCGGCCGCG (SEQ ID NO:11).

The oligonucleotides TCRIBamS and TCRIBstBS

(S oligonucleotides) were reverse-phase purified and had

their DMT protective group removed prior to use. The AS oligonucleotides were purified by reverse-phase, but

5' DMT groups were not removed. For each oligonucleotide pair, the sense (S) and antisence (AS) oligonucleotides were titrated against one another and, on the basis of these experiments, a 1:1 ratio (wt:wt) of S:AS oligonucleotides, for each oligonucleotide pair, was used to generate duplex substrate for reaction with Topo65.

The duplex substrates shown in Figures 3A and 3B were reacted overnight at room temperature with Topo65. The reaction mixtures contained 3  $\mu M$  Topo65,

10 μM duplex substrate, 10 mM Tris-HCl, pH 7.5, 100 mM KCl, 10 mM MgCl2, 1 mM EDTA, and 2 mM DTT. The reaction products were adjusted to 1 M NaCl or NOT and then analyzed by 10% SDS-PAGE and Coomassie Blue staining (lanes 4-7). Figure 4 is a photograph of the stained gel.

As a control, Topo65 was allowed to react with a duplex 22-mer suicide substrate. The suicide substrate contains a centrally positioned 5'-bridging phosphorothiolate. The sequences of the 22-mer oligonucleotides contained in the suicide substrate 22-mer duplex are (top strand) 5' AAAAAGACTTAGAAAAATTTTT 3' (SEQ ID NO:18 and (bottom strand) 5' AAAAATTTTTCTAAGTCTTTTT (SEQ ID NO:19). The position of the 5'-briding phosphorothiolate incorporated into the top oligonucleotide strand is between the sequences GACTT and AGAAAA. Topo65 quantitatively reacts with this suicide substrate, demonstrating that all of the enzyme is functional for cleavage.

Samples contained in each lane of the gel shown in Figure 4 are as follows: 1) Molecular weight markers; 2)

20 Topo65 reacted with suicide 22mer with salt; 3) Topo65 reacted with suicide 22-mer without salt; 4) Topo65 reacted with TCRIBamS/AS with salt; 5) Topo65 reacted with TCRIBamS/AS without salt; 6) Topo65 reacted with TCRIBstBS/AS with salt; 7) Topo65 reacted with TCRIBstBS/AS without salt; 7) Topo65 reacted with TCRIBstBS/AS without salt; and 8) Topo65, 2 μg.

Approximately 20% of the Topo65 can be trapped with SDS in covalent complex with DNA after cleavage of the TCRIBamAS/TCRIBamS or TCRIBstBS/TCRIBstBAS substrates. The SDS-PAGE gel revealed a shift in the mobility of Topo65 when

it is reacted with both TCRIBamS/AS and TCRIBstBS/AS, indicating that Topo65 can react at a cleavage site 2 to 4 base pairs from the 3' end of a duplex DNA. The mobility shift is equivalent to about 15 kDa, which is the expected molecular weight for a 22-mer attached to Topo65.

This experiment was repeated after removal of the DMT from the AS oligonucleotides by acid treatment, and an identical gel-shift was observed on reaction of "DMT off" oligonucleotide complexes with Topo65. However, no gel shift was observed if duplex generated from ratios of single strand oligonucleotides alone, for example, TCRIBamS, TCRIBstBS or TCRIBstBAS alone.

These results indicate that Topo65 cleaved at a 5'-phosphorothiolate moiety located two or four base pairs upstream from a blunt end.

### EXAMPLE III

# Topo65 Mediated Ligation of Oligonucleotide Substrates

This example shows Topo65 mediated ligation of oligonucleotide substrates following cleavage at the 5'-bridging phosphorothiolate located near the end of a DNA duplex.

The SDS-PAGE analysis depicted in Figure 4 demonstrates that topo I is capable of reacting with the oligo duplex substrates. In order to determine the fate of the substrates themselves, the duplex oligonucleotide substrates shown in Figure 5A were reacted with Topo65 and

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the oligonucleotide reaction products were examined by both native and denaturing PAGE (Figures 5A and 5B, respectively). Through this analysis, the efficient Topo65mediated generation of dimeric (about 50 bp) and even 5 trimeric (about 70 bp) DNA molecules was observed (Figure 5B, lanes 4 and 5).

Figure 5A depicts oligonucleotide substrates reacted with Topo65, labeled as Oligo pairs 1 through 6. Oligonucleotide substrates (pairs 1 to 3) were reacted with 10 Topo65, adjusted to 500 mM NaCl or NOT, and the resulting mixtures were digested with proteinase K in SDS prior to analysis by native PAGE (Figure 5B) or denaturing PAGE with urea (Figure 6B). Unreacted control oligonucleotide duplex samples are shown in lanes 14 and 15. Each gel contained 15 the following samples: (1) 20 bp DNA ladder, (2) oligonucleotide pair 1 with salt, (3), oligonucleotide pair 1 without salt, (4) oligonucleotide pair 2 with salt, (5) oligonucleotide pair 2 without salt, (6) oligonucleotide pair 4 with salt, (7) oligonucleotide pair 4 without salt, (8) oligonucleotide pair 5 with salt, (9) oligonucleotide pair 5 without salt, (10) oligonucleotide pair 3 with salt, (11) oligonucleotide pair 3 without salt, (12) oligonucleotide pair 6 with salt, (13) oligonucleotide pair 6 without salt, (14) 400 ng unreacted oligo pair 1, (15) 400 ng unreacted oligo pair 4.

Annealed "DMT ON" and "DMT OFF" oligo duplexes were reacted with Topo65 as described above. After overnight reaction at room temperature, 5M NaCl was added to a final concentration of 600mM and reaction continued for a further 2-3 hours at room temperature. Salt shifts the

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cleavage: ligation equilibrium of the top65 reaction towards The presence of 600mM NaCl thus results in an increased concentration of the DNA species produced by Topo65-mediated cleavage and relegation of duplex substrates 5 TCRIBamS/AS and TCRIBstBS/AS. The reaction was stopped with 0.1% SDS and then digested with proteinase-K (PK), for 30minutes at 37 degrees, to remove the Topo65 protein. product was analyzed on denaturing and native acrylamide gels. In all cases, DNA products of about 50bp and about 70bp were generated on reaction of Topo65 with TCRIBamS/AS (27mer:23mer) and TCRIBstBAS/AS (25mer:21mer). no differences in either size or quantity of the products generated using "DMT OFF" vs "DMT ON" oligo duplexes. The products observed using TCRIBstBS/AS as substrate were 15 consistently several base pairs smaller than those obtained with TCRIBamS/AS as substrate.

It should be noted that the Oligo pairs 1 and 3 were thought to contain DMT or phosphate blocking groups on the 5' end of each AS strand. The presence of such groups should have blocked topo I-mediated ligation events. 20 However, the observed lack of expected blocking is attributed to the apparent absence of 5' DMT or phosphate groups on the AS oligos, as determined by HPLC analysis). Hence, all three oligo pairs shown in Figure 5A were capable 25 of being cleaved by Topo65 at the 5' bridging phosphorothiolate moiety, and thenligated by Topo65 to a second oligo pair to generate a dimeric 50-mer.

# <u>Cloning and Sequence Analysis of the 50-mer Topo I Ligation</u> Product

The dimeric oligos of about 50 base pairs
resulting from Topo65 activity were gel purified. DNA
5 ligase was used to clone these fragments into an EcoRI
digested plasmid vector. Sequence analysis of clones
containing an insert demonstrated that in each case, Topo65
successfully ligated two oligonucleotides head to head.
This is consistent with the predicted product of Topo65
10 cleavage at the 5' phosphorothiolate moiety followed by
ligation of the 5' end of the AS strand to the 3' end of the
Topo65 cleaved 5' strand.

Once Topo65 has formed a covalent complex by cleaving a 5' phosphorothiolate-containing substrate, the activated complex can ligate directly to a blunt end of a second unreacted duplex, or can react with another covalent Topo65-oligonucleotide complex. The present assay does not discern between these two possibilities.

In both the case of the TCRIBamS/AS duplex and the TCRIBstBS/AS duplex, the sequence of the 50 mer product that is generated on reaction with Topo65 is that which is expected if cleavage and re-ligation occurs at the phosphorothiolate moiety. Figure 3C depicts the TCRIBamS/AS duplex after Topo65 cleavage. The GATC sequence removed on Topo65 cleavage yields a BamHI compatible 5' overhang. The sequence of the expected ligation product if two cleaved molecules are ligated at GATC overhang is shown in Figure 3D. This sequence is identical to the experimentally determined sequence of the product of topo-mediated ligation

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of the TCRIBams/AS duplex cloned into EcoRI sites of vector and sequenced using T3 and T7 primers. This sequencing data indicates that Topo65 cleaves the duplexes TCRIBams/AS and TCRIBstBAS at the 5' phosphorothiolate topo cleavage site to leave a 5' GATC or 5' GC overhang respectively.

The Topo65-mediated oligonucleotide linkage reaction with both BamHI (4 bases from the end ) and BstBI (2 bases from the end) sequences indicates that this human topoisomerase I-mediated ligation event can be achieved with 10 more than one sequence, in contrast to Vaccinia virus topoisomerase. Since it is well established that DNA ligase can not ligate a 5' OH to a 3' phosphate, the dimeric oligos which were cloned and subsequently sequenced could only have resulted from the activity of Topo65.

EXAMPLE IV

# Topo65 Mediated Circularization of 5'-Phosphorothiolate-Containing Plasmid DNA

This example shows that Topo65 can mediate circularization of linear plasmid DNA that has been adapted at both ends with oligonucleotide duplexes that contain a 5'-briding phosphorothiolate near the 5' end of the scissile strand.

To further examine the ability of Topo65 to

25 mediate ligation of DNAs containing 5'-briding
phosphorothicate near the 5' end of the scissile strand, a
vector was adapted with duplex oligonucleotides TCRIBamS/AS
described in Example II. Figure 6A depicts the linearized

and oligonucleotide adapted vectors. T4 DNA ligase was then used to adapt the vector ends with TCRIBamS oligonucleotide and TCRIBamAS oligonucleotides that contained or lacked a 5' phosphate. EcoR1-digested vector was incubated at 16°C 5 overnight with a 300-fold molar excess of oligonucleotides. The adapted vector was then incubated in the presence or absence of Topo65 in 10-fold molar excess, overnight at room temperature in a reaction buffer containing 10 mM Tris-HCl, pH 7.5, 100 mM KCl, 10 mM MgCl2, 1 mM EDTA and 2 mM DTT. After this incubation, 5 M NaCl was added to obtain a final concentration of 600 mM, and the mixture remained at room temperature for three hours. To this mixture, SDS was added to a final concentration of 0.1%, Proteinase-K was added to a final concentration of 100  $\mu g/ml$ , and the mixture was 15 incubated at 37°C for 30 minutes. The DNA was isolated by phenol/chloroform extraction, followed by ethanol precipitation. The isolated DNA was used to transform TOP10F' cells, which were then plated onto 2YT/Ampicillin agar.

The inserts obtained from 3-6 clones from each plate were sequenced and found to contain the head to head ligated oligonucleotides as predicted if Topo65 cleaves at the 5' phosphorothiolate moiety and then mediates ligation to an acceptor polynucleotide.

As depicted in Figure 6B, under conditions in which adaptored vector containing TCRIBamS and TCRIBamAS lacking a 5'-phosphate was incubated with Topo65, 4 of 6 clones were self-ligated adaptored pKSII and 2 of 6 clones had no insert. Under conditions in which adaptored vector containing TCRIBamS and TCRIBamAS having a 5'-phosphate was

incubated with Topo65, 2 of 6 clones were self-ligated adaptored pKSII and 1 of 6 clones had no insert, and 3 of 6 clones were re-circularized adaptored vector with deletion. Under conditions in which adaptored vector containing

5 TCRIBamS and TCRIBamAS lacking a 5'-phosphate was incubated in the absence of Topo65, 1 of 3 clones were re-circularized adaptored vector containing one adaptor, 1 of 3 clones were re-circularized adaptored vector with deletion, and 1 of 3 clones had no insert.

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### EXAMPLE V

### Purification of Topo65

This example shows a method of purifying Topo65 from baculovirus-infected sf9 insect cells.

Topo65 was prepared by infecting Sf9 insect cells 15 grown to about 2 X 106 cells/ml with a baculovirus containing a vector encoding Topo65 (Ac Topo65 viral stock passage 2). Insect cells were harvested 60 hours after infection by centrifugation at 1500rpm for 5 minutes at  $4^{\circ}$ C. Cells were then washed by combining cells in ~500ml PBS at 20 4°C and collected by centrifugation at 1500rpm for 5 minutes at 4°C. Topo65-expressing cells were then lysed by resuspending washed cells in 200ml lysis buffer (50 mM KCl, 10 mM Tris-HCl, pH 7.5, 2 mM MgCl2, 1% Triton X-100, 0.15 mg/ml PMSF, and 15 mM beta-mercaptoethanol). Nuclei were 25 collected by centrifugation at 2000rpm for 5 minutes at  $4^{\circ}$ C. Cytoplasmic supernatant was decanted. Collected nuclei were resuspended in 200ml resuspension buffer (50 mM KCl, 10 mM Tris-HCl, pH 7.5, 2 mM MgCl2, 0.15 mg/ml PMSF, and 15 mM beta-mercaptoethanol) and then centrifuged at 2000rpm for

5 minutes at 4°C. Buffer was then removed, and nuclei were washed a second time.

Washed nuclei were resuspended in 40ml resuspension buffer containing fresh PMSF and beta-mercaptoethanol containing 0.8 ml 0.5M EDTA. Nuclei were then lysed by addition of 45ml 2X nuclear lysis buffer (80 mM Tris-HCl, pH 7.5, 2 M NaCl, 20% glycerol, 2 mM EDTA) followed by shaking of mixture. DNA was precipitated from the mixture by dropwise addition 50ml PEG solution (18% PEG-8000, 1 M NaCl, 10% Glycerol) to mixture stirring at 4°C. The mixture was then allowed to stir at 4°C for 5 hours.

To remove precipitated DNA from the nuclei mixture, centrifugation was performed at 8000rpm for 10 minutes at  $4^{\circ}$ C, followed by removal of supernatant. 15 PEG supernatant was dialyzed against 4 liters dialysis buffer (250 mM KPO4, 1 mM EDTA, 0.1 mM PMSF, and 5 mM betamercaptoethanol) overnight at  $4^{\circ}\text{C}$ . To clarify dialyzed PEG supernatant, centrifugation at was performed at 8000rpm for 20 minutes at 4°C. Topo65 was purified from clarified PEG supernatant by phenyl sepharose chromatography. A column 20 (approximately 10 cm) was packed with Phenyl Sepharose CL-4B (Pharmacia cat. # 17-0810-01, Pharmacia, Piscataway, NJ) and equilibrated with dialysis buffer. Clarified PEG supernatant was loaded onto a phenyl sepharose column by gravity and fractions were collected on ice. The phenyl sepharose flow through (~200ml) was diluted with 200ml of POROS HS buffer A (10 mM Tris-HCl, pH 7.5, 1 mM EDTA, 5 mM beta-mercaptoethanol, 0.1 mM PMSF).

POROS 20 HS Chromatography (Perseptive Biosystems matrix, cat. # 1-3329-06) chromatography (3.32 ml) was performed on Phenyl Sepharose flow through. The column was washed with buffer B (POROS HS buffer A plus 1 M KCl) and 5 then equilibrated column with Buffer A. The phenyl sepharose flowthrough over the equilibrated column at a rate of 2 ml/minute. When all flowthrough was been passed over the column, re-equilibration was performed with 4 column volumes of Buffer A. Protein was eluted from the column with a 15-800mM KCl gradient. SDS-PAGE was performed using 5-10ul aliquots of each fraction collected. Fractions containing Topo65 protein (fractions 25-30) were pooled. Topo65 was further purified by MonoQ chromatography of pooled fractions from POROS 20 column. Pooled fractions 15 were diluted with 2 volumes of water and loaded onto a MonoQ column equilibrated with MonoQ Buffer A (25 mM KPO4, pH 7.4, 1 mM EDTA, 1 mM beta-mercaptoethanol, 1 mM PMSF), followed by elution with MonoQ buffer B (1 M KPO4, pH 7.4, 1 mM EDTA, 1 mM beta-mercaptoethanol, 1 mM PMSF).

Further purification of Topo65 was performed by MonoS Chromatography on a MonoS column equilibrated with MonoS buffer A (25 mM KPO4, pH 7.4, 1 mM EDTA, 1 mM betamercaptoethanol, 1 mM PMSF). Pooled Topo65 eluate from MonoQ chromatography was loaded onto a MonoS column followed by elution with a 7.5-40% gradient of MonoS buffer B (1 M KPO4, pH 7.4, 1 mM EDTA, 1 mM beta-mercaptoethanol, 1 mM PMSF) over 20 column volumes. SDS-PAGE was performed to detect Topo65 and fractions containing Topo65 protein (fractions 9-13) were pooled. Pooled Fractions from MonoS chromatography were dialyzed overnight against final dialysis buffer at 4°C (10 mM Tris-HCL, pH 7.5, 1 mM EDTA,

and 1 mM DTT). The final concentration of dialyzed material was adjusted to 5 mM DTT and Topo65 concentration was determined by Bradford Assay. The procedure yielded 6 ml of Topo65 at a concentration of 2 mg/ml.

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### EXAMPLE VI

## Synthesis of Polynucleotide-5' Phosphorothiolates

This example describes the synthesis of two polynucleotide 5'-phosphorothiolates.

Figure 7A shows a schematic representation of the chemical synthesis of the 5'-phosphorothiolate nucleic acid 5'-S-dimethoxytrityl-3'-O-cyanoethyl-NN-diisopropyl-phosphoramidite-2'-deoxy-N-benzoyl-cytosine. Preparation of the compound was performed as follows:

N-benzoyl-2'-deoxy cytosine (3.13 mmoles) was dissolved in anhydrous tetrahydrofuran (THF) at 70°C and allowed to cool to 25°C. Triphenyl phosphine (3.75 mmoles), diethyl azodicarboxylate (3.29 mmoles), and thiolacetic acid (3.44 mmoles) were added sequentially. The reaction was stirred at 25°C for 6 hours and quenched by adding 5ml anhydrous methanol, and dried in vacuo. The residue was resuspended in ethyl acetate, and extracted with 3 volumes 5% sodium bicarbonate. Organic layer was dried in vacuo and the product purified silica gel chromatography (5% methanol in dichloromethane).

5'-thio-acetate derivative was dissolved in 20 ml anhydrous methanol and 10ml anhydrous pyridine was added. 6

ml of 2N NaOH in methanol was added, and the reaction was then stirred at room temp for 5 minutes. The reaction was cooled to 0°C, and 975 ul of concentrated HCL was added followed by 100 ml of 5% phosphate buffer pH 7.0. The aqueous phase was extracted 2X with 100ml ethyl acetate. The organic phase was dried in vacuo and the 5'-deoxy-5'-sulfhydryl nucleoside was purified by silica gel chromatography (10% methanol in dichloromethane). All solvents were extensively degassed and purged with argon.

- 5'-deoxy-5'-sulfhydryl-2'-deoxy-N-benzoylcytosine (2 mmoles) was dissolved in anhydrous pyridine and dimethoxy-trityl chloride (2.2 mmoles) was added under argon. The reaction was stirred at room temperature for 1 hour, and then quenched by adding 100 ul of

  15 beta-mercaptoethanol. The reaction was dried in vacuo, resuspended in dichloromethane, and purified by silica gel chromatography (5% methanol, 0.5% beta-mercaptoethanol in dichloromethane).
- 5'-S-dimethoxytrityl-2'-deoxy-N-benzoyl
  cytosine (1 mmole) was dissolved in anhydrous acetonitrile and NN-diisopropyl-ethylamine (1.2 mmoles) and 2-cyanoethyl diisopropyl-chlorophosphoramidite (1.1 mmoles) were added sequentially at room temp. The reaction was stirred for 60 minutes and quenched with 100 ul of anhydrous methanol.

  The mixture was dried in vacuo and the product purified by silica gel chromatography (45% ethyl acetate, 25% dichloromethane, 10% triethylamine).

Figure 7B shows a schematic representation of the chemical synthesis of 5'-S-dimethoxytrityl-3'-O-

cyanoethyl-NN-diisopropyl-phosphoramidite-2'-deoxy-N-isobuty ryl quanosine. The compound was prepared as follows:

N-benzoyl-2'-deoxy cytosine (3.13 mmoles) was dissolved in anhydrous dimethyl formamide (DMF). Triphenyl phosphine (3.75 mmoles), diethyl azodicarboxylate (3.29 mmoles), and thiolacetic acid (3.44 mmoles) were added sequentially. The reaction was stirred at 25°C for 6 hours and quenched by adding 5 ml anhydrous methanol. Two volumes of water were added and mixture cooled to -20°C. The precipitate was collected by filtration, resuspended in ethyl acetate, and extracted with 3 volumes 5% sodium bicarbonate. The resulting organic layer was dried in vacuo and the product purified by silica gel chromatography (10% methanol in dichloromethane).

- anhydrous methanol and 10 ml anhydrous pyridine was added.
  6 ml of 2N NaOH in methanol was added, and the reaction was then stirred at room temp for 5 minutes. The reaction was cooled to 0°C, and 975 ul of concentrated HCL was added followed by 100 ml of 5% phosphate buffer pH 7.0. The aqueous phase was extracted 2X with 100 ml ethyl acetate. The organic phase was dried in vacuo and the 5'-deoxy-5'-sulfhydryl nucleoside was purified by silica gel chromatography (10% methanol in dichloromethane).

  Note: all solvents were extensively degassed and purged
- 25 Note: all solvents were extensively degassed and purged with argon.
  - 5'-deoxy-5'-sulfhydryl-2'-deoxy-N-isobutyryl quanosine (2 mmoles) was dissolved in anhydrous pyridine and

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dimethoxy-trityl chloride (2.2 mmoles) was added under argon. The reaction was stirred at room temperature for 1 hour, and then quenched by adding 100 ul of beta-mercaptoethanol. The reaction was dried in vacuo, resuspended in dichloromethane, and purified by silica gel chromatography (5% methanol, 0.5% beta-mercaptoethanol in dichloromethane).

5'-S-dimethoxytrityl-2'-deoxy-N-isobutyryl guanosine (1 mmole) was dissolved in anhydrous acetonitrile

10 and NN-diisopropyl-ethylamine (1.2 mmoles) and 2-cyanoethyl diisopropyl-chlorophosphoramidite (1.1 mmoles) were added sequentially at room temp. The reaction was stirred for 60 minutes and quenched with 100 ul of anhydrous methanol. The mixture was dried in vacuo and the product purified by silica gel chromatography (45% ethyl acetate, 25% dichloromethane, 10% triethylamine).

The phosphoramidites were added to the growing oligonucleotide chain using an ABI 392 DNA synthesizer using standard reagents with the exception of the 5'-S-DMT deprotection step.

The 5'-O-DMT protecting group was removed using 2-3% trichloroacetic acid in dichloromethane (standard protocol). Heavy metals (e.g. silver nitrate or mercuric chloride) have been used to remove the 5'-S-Trityl or 5'-S-DMT group. It was observed that 7.5% trichloroacetic acid, 1% betamercaptoethanol in dichloromethane, followed by extensive washing with degassed (argon purged) anhydrous acetonitrile was efficient for removing the DMT and exposing the 5'-sulfhydryl for coupling. To form the

phosphorothiolate linkage, the coupling time was increased to 300 minutes.

### EXAMPLE VII

## Synthesis of a Polynucleotide-3' Phosphorothiolate

This Example shows an exemplary protocol for one step synthesis of a polynucleotide-3' phosphorothiolate.

A polynucleotide-3' phosphorothiolate (3'-SNP) was prepared by incubating 3'-monophosphorothiolate polynucleotide with 4'-iodo-nitrophenyl, as shown in 10 Figure 8. Oligonucleotides containing a 3'monophosphorothioate were synthesized using 3'-phosphate CPG (Glen Research, Sterling, VA) and standard DNA phosphoramidites on an ABI 392 DNA synthesizer. The first coupling step was sulfurized, rather than oxidized, so that 15 polynucleotide containing a 3'-monophosphorothioate, rather than a 3'-phosphate, was produced following deprotection. The purified oligonucleotide was incubated with saturating concentrations of 4'-iodo-nitrophenyl (Aldrich, St. Louis, Missouri) in 50% water, 50% acetonitrile, 50 mM MES, pH 6.0, 1 mM  $MgCl_2$ , overnight at 37°C with vigorous shaking. 20 Figure 8, the starting material 3'-monophosphorothioate polynucleotide is shown on the left, the product 3'-SNP polynucleotide is shown on the right, while the activator, 4-iodo-nitrophenyl, is pictured in the center.

After incubation, two volumes of water were added to the reaction mixture, and the mixture was clarified by centrifugation. The solution was then extracted three times with ethyl acetate, and the polynucleotide-3'

phosphorothicate is precipitated from the aqueous phase by adding one tenth volume of 3M sodium acetate and three volumes of absolute ethanol.

Thus, a polynucleotide-3' phosphorothiolate was prepared by reacting a polynucleotide-3' phosphorothiolate precursor, 3'-monophosphorothioate polynucleotide, with an activator, 4-iodo-nitrophenyl, using commercially available reagents.

Throughout this application various publications

10 have been referenced within parentheses. The disclosures of these publications in their entireties are hereby incorporated by reference in this application in order to more fully describe the state of the art to which this invention pertains.

15 Although the invention has been described with reference to the disclosed embodiments, those skilled in the art will readily appreciate that the specific experiments detailed are only illustrative of the invention. It should be understood that various modifications can be made without 20 departing from the spirit of the invention.